=> fil hcaplus FILE 'HCAPLUS' ENTERED AT 07:05:26 ON 23 JUL 2003 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2003 AMERICAN CHEMICAL SOCIETY (ACS)

Jan Delaval Reference Librarian Biotechnology & Chemical Library CM1 1E07 – 703-308-4498 jan.delaval@uspto.gov

Copyright of the articles to which records in this database refer is held by the publishers listed in the PUBLISHER (PB) field (available for records published or updated in Chemical Abstracts after December 26, 1996), unless otherwise indicated in the original publications. The CA Lexicon is the copyrighted intellectual property of the the American Chemical Society and is provided to assist you in searching databases on STN. Any dissemination, distribution, copying, or storing of this information, without the prior written consent of CAS, is strictly prohibited.

FILE COVERS 1907 - 23 Jul 2003 VOL 139 ISS 4 FILE LAST UPDATED: 22 Jul 2003 (20030722/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> d all tot

```
L66 ANSWER 1 OF 10 HCAPLUS COPYRIGHT 2003 ACS on STN
```

AN 2002:275805 HCAPLUS

DN 136:314972

- TI Compositions and methods for the transport of biologically active agents across cellular barriers
- IN Houston, Lou L.; Sheridan, Philip J.; Hawley, Stephen; Glynn, Jacqueline
 M.; Chapin, Steven; Basu, Amaresh
- PA Arizeke Pharmaceuticals, Inc., USA
- SO PCT Int. Appl., 378 pp. CODEN: PIXXD2
- DT Patent
- LA English
- IC ICM A61K038-00
- CC 63-5 (Pharmaceuticals)

Section cross-reference(s): 1, 2, 9, 15

FAN.CNT 1

r MN.	-					KIND		DATE		APPLICATION NO.					DATE			
PI		2002028408								WO 2001-US30832 20011002								
		W:	AE, CO, GM, LS, PT, US, GH, DE,	AG, CR, HR, LT, RO, UZ, GM, DK,	AL, CU, HU, LU, RU, VN, KE, ES,	AM, CZ, ID, LV, SD, YU, LS, FI,	AT, DE, IL, MA, SE, ZA, MW, FR,	AU, DK, IN, MD, SG, ZW, MZ, GB,	DM, IS, MG, SI, AM, SD, GR,	DZ, JP, MK, SK, AZ, SL, IE,	EC, KE, MN, SL, BY, SZ, IT,	EE, KG, MW, TJ, KG, TZ, LU,	ES, KP, MX, TM, KZ, UG, MC,	FI, KR, MZ, TR, MD, ZW, NL,	GB, KZ, NO, TT, RU, AT, PT,	GD, LC, NZ, TZ, TJ, BE, SE,	GE, LK, PH, UA, TM CH,	GH, LR, PL, UG,
PRAI	US US US	20010 13247 R: 20000 20000 20000 2001	0964: 778 AT, IE, -237: -248:	94 BE, SI, 929P 478P 819P	A: CH, LT, P P	5 2 DE, LV,	2003 DK, FI, 2000 2000 2000	0415 0709 ES, RO, 1002 1113	FR, MK,	AI El GB,	U 200 P 200 GR,	01-9 01-9 IT,	6494 7736	3	2001: 2001:	1002 1002		PT,

20011002 WO 2001-US30832 W Disclosed herein are complexes and compds. that pass through cellular AB barriers to deliver compds. into, through and out of cells, and methods of producing and using such complexes and compds. The complexes and compds. of the invention comprise a biol. active portion and a targeting element directed to a ligand that confers transcellular, transcytotic or paracellular transporting properties to an agent specifically bound to the ligand, with the proviso that the targeting element is not an antibody. Also disclosed are complexes and compds. that comprise two or more targeting elements directed to a ligand that confers transcellular, transcytotic or paracellular transporting properties to an agent specifically bound to the ligand. Preferred ligands include but are not limited to the stalk of plgR, a plgR domain, an amino acid sequence that is conserved among plgR's from different animals, and one of several regions of pIgR defined herein. drug targeting ${\tt pIgR}$ cell delivery endocytosis ST Proteins ΙT RL: BSU (Biological study, unclassified); PRP (Properties); BIOL (Biological study) (AP-1 (adaptor-related protein complex 1), peptides; compns. and methods for the transport of biol. active agents across cellular barriers) Proteins ΙT RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (adaptor; compns. and methods for the transport of biol. active agents across cellular barriers) TT Diagnosis (agents; compns. and methods for the transport of biol. active agents across cellular barriers) IT Endocytosis (apical; compns. and methods for the transport of biol. active agents across cellular barriers) ΙT Diagnosis Drug delivery systems Endocytosis Human Macaca fascicularis ·Macaca mulatta Molecular cloning Peptidomimetics Pharmacokinetics Protein sequences Signal transduction, biological Test kits Transformation, genetic cDNA sequences (compns. and methods for the transport of biol. active agents across cellular barriers) Fusion proteins (chimeric proteins) IT RL: BPN (Biosynthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (compns. and methods for the transport of biol. active agents across cellular barriers) ΙT Nucleic acids RL: PEP (Physical, engineering or chemical process); PYP (Physical process); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses) (compns. and methods for the transport of biol. active agents across cellular barriers) Antibodies ΙT

Antigens

Blood-coagulation factors

Carbohydrates, biological studies Enzymes, biological studies Growth factors, animal Hormones, animal, biological studies Interleukin 2 Interleukin 4 Interleukins Ligands Lipids, biological studies Receptors Transcription factors Transport proteins RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (compns. and methods for the transport of biol. active agents across cellular barriers) IT Amniotic fluid Blood Lymph (delivery to; compns. and methods for the transport of biol. active agents across cellular barriers) TT Body fluid (interstitial, delivery to; compns. and methods for the transport of biol. active agents across cellular barriers) Biological transport IT. (intracellular; compns. and methods for the transport of biol. active agents across cellular barriers) ΙT Drug delivery systems (liposomes; compns. and methods for the transport of biol. active agents across cellular barriers) TΤ Bladder Digestive tract Eye Lung Nose Uterus Vagina (lumen; compns. and methods for the transport of biol. active agents across cellular barriers) TΤ Proteins RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (nucleic acid-binding; compns. and methods for the transport of biol. active agents across cellular barriers) ΙT Immunoglobulin receptors RL: BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (pIgR (polymeric Ig receptor); compns. and methods for the transport of biol. active agents across cellular barriers) TΨ Calmodulins RL: BSU (Biological study, unclassified); PRP (Properties); BIOL (Biological study) (peptides; compns. and methods for the transport of biol. active agents across cellular barriers) Proteins TT RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (scaffolding; compns. and methods for the transport of biol. active agents across cellular barriers) TΤ Drug delivery systems (targeted; compns. and methods for the transport of biol. active agents across cellular barriers) 407584-46-7 ፐጥ 407584-44-5 407584-45-6 407584-47-8 407584-48-9 407584-49-0 407584-50-3 407584-51-4 RL: PRP (Properties); THU (Therapeutic use); BIOL (Biological study); USES

```
(Uses)
        (compns. and methods for the transport of biol. active agents across
        cellular barriers)
     7440-05-3, Palladium, biological studies
                                               7440-06-4, Platinum, biological
ΙT
              7440-48-4, Cobalt, biological studies
                                                     7440-66-6, Zinc,
                          9001-92-7, Proteinase 9002-60-2, Corticotropin,
     biological studies
                          9002-72-6, Somatotropin 9004-10-8, Insulin,
     biological studies
                                                  372092-80-3, Protein kinase
                          9013-05-2, Phosphatase
     biological studies
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (compns. and methods for the transport of biol. active agents across
        cellular barriers)
                                 409334-92-5
                                               409334-93-6
IT
     245322-12-7
                  250649-36-6
                                                             409334-94-7
                                               409334-99-2
     409334-95-8
                   409334-97-0
                                 409334-98-1
                                                             409335-01-9
     409335-03-1
                  409335-05-3
                                 409412-43-7
                                               409412-44-8
                                                             409412-45-9
     409412-46-0
                  409412-47-1
                                 409412-48-2
                                               409412-49-3
                                                             409412-50-6
                                               409412-54-0
     409412-51-7
                  409412-52-8
                                 409412-53-9
                                                             409412-55-1
     409412-56-2
                  409412-57-3
                                 409412-58-4
                                               409412-59-5
                                                             409412-60-8
                                               409412-64-2
                                                             409412-65-3
     409412-61-9
                  409412-62-0
                                 409412-63-1
                                               409412-69-7
                                                             409412-70-0
     409412-66-4
                  409412-67-5
                                 409412-68-6
                                               409412-74-4
                                                             409412-75-5
     409412-71-1
                  409412-72-2
                                 409412-73-3
                                               409412-79-9
                                                             409412-80-2
     409412-76-6
                  409412-77-7
                                 409412-78-8
     409412-81-3
                  409412-82-4
                                 409412-83-5
                                               409412-84-6
                                                             409412-85-7
                                               409412-89-1
                                                             409412-90-4
     409412-86-8
                  409412-87-9
                                 409412-88-0
                                               409412-94-8
                                                             409412-95-9
     409412-91-5
                  409412-92-6
                                 409412-93-7
                                               409412-99-3
                                                             409413-00-9
     409412-96-0
                  409412-97-1
                                 409412-98-2
                                               409413-04-3
                                                             409413-05-4
     409413-01-0
                  409413-02-1
                                 409413-03-2
     409413-06-5
                  409413-07-6
                                 409413-08-7
                                               409413-09-8
                                                             409413-10-1
                                                             409413-15-6
     409413-11-2
                  409413-12-3
                                 409413-13-4
                                               409413-14-5
                                 409413-18-9
                                               409413-19-0
                                                             409413-20-3
     409413-16-7
                  409413-17-8
                                 409413-23-6
                                               409413-24-7
                                                             409413-25-8
     409413-21-4
                  409413-22-5
                                               409413-29-2
                                                             409413-30-5
                  409413-27-0
                                 409413-28-1
     409413-26-9
                   409413-32-7
                                 409413-33-8
     409413-31-6
     RL: PRP (Properties)
        (unclaimed sequence; compns. and methods for the transport of biol.
        active agents across cellular barriers)
    ANSWER 2 OF 10 HCAPLUS COPYRIGHT 2003 ACS on STN
L66
AN
     2001:823032 HCAPLUS
DN
     137:31596
ΤI
     Immunobiology of secretory IgA antibodies
ΑU
     Department of Dentistry, Nihon University, Japan
CS
     Nenmaku Men'eki (2001), 113-133, 294. Editor(s): Kiyono, Hiroshi;
SO
     Ishikawa, Hiromichi; Nagura, Hiroshi. Publisher: Nakayama Shoten, Tokyo,
     Japan.
     CODEN: 69BZKP
DΤ
     Conference; General Review
LA
     Japanese
CC
     15-0 (Immunochemistry)
     A review discussing structures and functions of structural components of
AB
     secretory IgA. The structural components of secretory
     IgA are IgA, J chain, and secretory component polymeric
     Ig receptor. The review includes structure and function
     of secretory and membrane IgA; secretory IgA
     expression; structure and function of secretory component
     polymeric Ig receptor; genesis of
     polymeric Ig receptor; role of cytokines in
     polymeric Ig receptor expression; and
     structure and function of J chain.
ST
     review secretion IgA Ig
ΙT
     Immunoglobulins
     RL: BSU (Biological study, unclassified); PRP (Properties); BIOL
     (Biological study)
```

```
(A, secretory; secretory IgA structural components
        IgA, J chain, and secretory component)
IT
     Immunoglobulins
     RL: BSU (Biological study, unclassified); PRP (Properties); BIOL
     (Biological study)
        (A; secretory IgA structural components IgA, J chain, and
        secretory component)
     Immunoglobulins
ΙT
     RL: BSU (Biological study, unclassified); PRP (Properties); BIOL
     (Biological study)
        (fragments, J-chain; secretory IgA structural
        components IgA, J chain, and secretory component)
ΙT
     Immunoglobulins
     RL: BSU (Biological study, unclassified); PRP (Properties); BIOL
     (Biological study)
        (secretory component; secretory IgA structural
        components IgA, J chain, and secretory component)
     ANSWER 3 OF 10 HCAPLUS COPYRIGHT 2003 ACS on STN
L66
     2001:730836 HCAPLUS
ΑN
DN
     135:287529
     Ligands directed to the non-secretory component, non-
ΤI
     stalk region of plgR and methods of use thereof
     Mostov, Keith E.; Chapin, Steven J.;
ΙN
     Richman-Eisenstat, Janice
PΑ
     Regents of the University of California, USA
SO
     PCT Int. Appl., 102 pp.
     CODEN: PIXXD2
DT
     Patent
     English
LA
TC
     ICM C07K016-28
     ICS A61K039-395; A61K048-00; A61K038-00; A61K031-00; A61K031-7088;
          A61K047-48; C07K019-00; A61P011-00; C07K014-705
CC
     15-3 (Immunochemistry)
     Section cross-reference(s): 1, 3, 8, 63
FAN.CNT 1
     PATENT NO.
                       KIND DATE
                                             APPLICATION NO.
                                                               DATE
                             -----
                                             -----
     -----
                      ____
                     A2
     WO 2001072846
                             20011004
                                             WO 2001-US9699
                                                               20010326 <--
PΙ
     WO 2001072846
                       A3
                             20020404
            AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CO, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM,
             HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS,
             LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO,
             RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
             BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                                             20010326 <--
                       A1
                             20020801
                                            US 2001-818247
     US 2002102657
                             20030102
                                                              20010326 <--
                                             EP 2001-926437
     EP 1268555
                        Α2
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
PRAI US 2000-192197P
                             20000327
                      P
                           20000327
     US 2000-192198P
                       Ρ
     WO 2001-US9699
                       W
                             20010326 <--
     The invention provides compns. and methods for specific binding to a
AB
     region of the polymeric Ig receptor (
     pIgR) of a cell with the provisos that the ligand does
     not substantially bind to the most abundant form of the secretory
     component (SC) of \operatorname{\textbf{pIgR}} present in an organ of interest of an
     animal of interest under physiol. conditions, and does not bind to the
     plgR stalk. In some embodiments, the ligand
```

decreases cleavage of SC from the stalk by at least one-third. The ligands and methods of the invention can be used with both birds and mammals. In more preferred embodiments, the animal is a mammal. In the most preferred embodiment, the animal is a human. The ligand may be targeted into the cell or may undergo retrograde transcytosis and release at the basolateral side of the cell, and may comprise a biol. active compn. STpolymeric Ig receptor secretory component ligand; antibody polymeric Ig receptor secretory component; epithelial cell drug delivery antibody pIgR ΙT Disulfide group (antibody fragment stabilization; ligands or antibodies directed to the non-secretory component, non-stalk region of polymeric Ig receptor plgR for drug targeting or delivery) Cell membrane TΤ (apical, epithelial; ligands or antibodies directed to the non-secretory component, non-stalk region of polymeric Ig receptor pIgR for drug targeting or delivery) Cell membrane ΙT (basolateral; ligands or antibodies directed to the non-secretory component, non-stalk region of polymeric Ig receptor pIgR for drug targeting or delivery) Organic compounds, biological studies RL: BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (biol.; ligands or antibodies directed to the nonsecretory component, non-stalk region of polymeric Ig receptor pIgR for drug targeting or delivery) ΙT Antibodies RL: BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (conjugates; ligands or antibodies directed to the non-secretory component, non-stalk region of polymeric Ig receptor pIgR for drug targeting or delivery) TT Immunoglobulins RL: BPN (Biosynthetic preparation); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (fragments; ligands or antibodies directed to the non-secretory component, non-stalk region of polymeric Ig receptor pIgR for drug targeting or delivery) TT Animal (human; ligands or antibodies directed to the nonsecretory component, non-stalk region of polymeric Ig receptor pIgR for drug targeting or delivery) IT Drug delivery systems (immunoconjugates; ligands or antibodies directed to the non-secretory component, non-stalk region of polymeric Ig receptor pIgR for drug targeting or delivery) IT Biological transport (internalization; ligands or antibodies directed to the non-secretory component, non-stalk region of polymeric Ig receptor pIgR for drug targeting or delivery)

```
Intestine
ΙT
        (large; ligands or antibodies directed to the non-
        secretory component, non-stalk region of
        polymeric Ig receptor pIgR for
        drug targeting or delivery)
ΙT
     Animal cell
     Anti-infective agents
     Anti-inflammatory agents
     Antibiotics
     Biliary tract
     Bird (Aves)
     Cat (Felis catus)
     Cattle
     Dog (Canis familiaris)
     Epithelium
     Epitopes
     Horse (Equus caballus)
     Lacrimal gland
     Liver
    Lung
     Mammal (Mammalia)
     Mammary gland
     Molecular cloning
     Nose
       Organ, animal
     Peptidomimetics
     Protein sequences
     Salivary gland
     Sheep
     Stomach
     Swine
     Uterus
     Vaqina
        (ligands or antibodies directed to the non-
        secretory component, non-stalk region of
        polymeric Ig receptor pIgR for
        drug targeting or delivery)
TT
     Antibodies
     Fusion proteins (chimeric proteins)
     RL: BPN (Biosynthetic preparation); BSU (Biological study, unclassified);
     THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
        (ligands or antibodies directed to the non-
        secretory component, non-stalk region of
        polymeric Ig receptor pIgR for
        drug targeting or delivery)
ΙT
     Antisense oligonucleotides
     CFTR (cystic fibrosis transmembrane conductance regulator)
     Carbohydrates, biological studies
       Ligands
     Lipids, biological studies
     Nucleic acids
     Proteins, general, biological studies
     Radionuclides, biological studies
     RL: BSU (Biological study, unclassified); THU (Therapeutic use); BIOL
     (Biological study); USES (Uses)
        (ligands or antibodies directed to the non-
        secretory component, non-stalk region of
        polymeric Ig receptor pIgR for
        drug targeting or delivery)
     Animal cell
IΤ
        (mammalian; ligands or antibodies directed to the
```

```
non-secretory component, non-stalk region of
        polymeric Ig receptor pIgR for
        drug targeting or delivery)
IT
     Drug delivery systems
        (mucosal; ligands or antibodies directed to the
        non-secretory component, non-stalk region of
        polymeric Ig receptor pIgR for
        drug targeting or delivery)
IT
     Immunoglobulin receptors
     RL: BPN (Biosynthetic preparation); BPR (Biological process); BSU
     (Biological study, unclassified); PRP (Properties); THU (Therapeutic use);
     BIOL (Biological study); PREP (Preparation); PROC (Process); USES (Uses)
        (polymeric Ig; ligands or
        antibodies directed to the non-secretory component,
        non-stalk region of polymeric Ig
        receptor pIgR for drug targeting or delivery)
IT
     Transcytosis
        (receptor-mediated; ligands or antibodies
        directed to the non-secretory component, non-stalk
        region of polymeric Ig receptor
        pIgR for drug targeting or delivery)
IT
     Immunoglobulin receptors
     RL: BPN (Biosynthetic preparation); BPR (Biological process); BSU
     (Biological study, unclassified); PRP (Properties); THU (Therapeutic use);
     BIOL (Biological study); PREP (Preparation); PROC (Process); USES (Uses)
        (secretory component; ligands or antibodies
        directed to the non-secretory component, non-stalk
        region of polymeric Ig receptor
       pIgR for drug targeting or delivery)
ΙT
     Body, anatomical
        (sinus; ligands or antibodies directed to the non-
        secretory component, non-stalk region of
        polymeric Ig receptor pIgR for
        drug targeting or delivery)
TΤ
     Intestine
     Molecules
        (small; ligands or antibodies directed to the non-
        secretory component, non-stalk region of
        polymeric Ig receptor pIgR for
        drug targeting or delivery)
                                 365241-70-9
                                               365295-56-3
TΤ
     365241-68-5
                   365241-69-6
     RL: PRP (Properties)
        (unclaimed sequence; ligands directed to the non-
        secretory component, non-stalk region of plgR
        and methods of use thereof)
    ANSWER 4 OF 10 HCAPLUS COPYRIGHT 2003 ACS on STN
L66
ΑN
     2000:756900 HCAPLUS
DN
     133:331775
ΤI
     Protein transport assays using IR fluorescent labeled ligands
     Mostov, Keith; Altschuler, Yoram
IN
     Regents of the University of California, USA
PA
SO
     PCT Int. Appl., 25 pp.
     CODEN: PIXXD2
DT
     Patent
LA
     English
     ICM C12Q001-00
IC
         C12Q001-02; C12Q001-04; C12Q001-32; G01N033-00; G01N033-53;
          C07H019-20
CC
     9-5 (Biochemical Methods)
     Section cross-reference(s): 1, 6, 15
FAN.CNT 1
     PATENT NO.
                      KIND DATE
                                           APPLICATION NO.
```

```
WO 2000063418
                            20001026
                                           WO 2000-US10173
                                                            20000414 <--
                      A 1
PΤ
        W: AU, CA, JP
        RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
             PT, SE
                            19990415 <--
PRAI US 1999-292274
                       Α
    MARPAT 133:331775
     The invention provides methods and compns. for quant. detecting
AΒ
     ligand movement across a biol. membrane. The general method
     comprises the steps of (a) contacting a ligand comprising an
     assay-compatible IR fluorescent label with a receptor under conditions
     wherein the receptor transports an amt. of the ligand across a
     biol. membrane; and (b) quant. detecting fluorescence as an indicator of
     the amt. of the ligand transported across the membrane. IgA
     labeled with NN382 or Cy5.5 was used to examine drugs affecting IgA
     transport in MDCK cells transfected with cDNA for rabbit pIgR.
ST
    protein transport assay IR fluorescent labeled ligand; IgA
     transport drug screening
     Immunoglobulins
ΙT
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (A, detection of drugs affecting transport of, across epithelial cell
        membranes; protein transport assays using IR fluorescent labeled
        ligands)
     Fluorescent indicators
IT
     Fluorescent probes
        (IR fluorescent ligands; protein transport assays using IR
        fluorescent labeled ligands)
     Animal cell line
ΙT
        (MDCK, in detection of drugs affecting transport of IgA across
        epithelial cell membranes; protein transport assays using IR
        fluorescent labeled ligands)
ΙT
     Fluorescent dyes
        (as labels; protein transport assays using IR fluorescent labeled
        ligands)
     Immunoglobulins
ΙT
     RL: ARG (Analytical reagent use); BPR (Biological process); BSU
     (Biological study, unclassified); ANST (Analytical study); BIOL
     (Biological study); PROC (Process); USES (Uses)
        (conjugates, A, with fluorescent labels; protein transport assays using
        IR fluorescent labeled ligands)
     Transferrins
IΤ
     RL: ARG (Analytical reagent use); BPR (Biological process); BSU
     (Biological study, unclassified); ANST (Analytical study); BIOL
     (Biological study); PROC (Process); USES (Uses)
        (conjugates, with fluorescent labels; protein transport assays using IR
        fluorescent labeled ligands)
TΤ
     Ligands
     RL: ARG (Analytical reagent use); BPR (Biological process); BSU
     (Biological study, unclassified); ANST (Analytical study); BIOL
     (Biological study); PROC (Process); USES (Uses)
        (contg. IR fluorescent labels; protein transport assays using IR
        fluorescent labeled ligands)
TΤ
     Biological transport
        (intracellular; protein transport assays using IR fluorescent labeled
        ligands)
     Proteins, specific or class
TT
     RL: ARG (Analytical reagent use); BPR (Biological process); BSU
     (Biological study, unclassified); ANST (Analytical study); BIOL
     (Biological study); PROC (Process); USES (Uses)
        (labeled, contg. IR fluorescent labels; protein transport assays using
        IR fluorescent labeled ligands)
TΤ
     Endosome
```

```
(membrane; protein transport assays using IR fluorescent labeled
        ligands)
    Transport proteins
TT
    RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (norepinephrine-transporting, noradrenaline transport through; protein
        transport assays using IR fluorescent labeled ligands)
    Transport proteins
IT
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (peptide-transporting, PEPT1; protein transport assays using IR
        fluorescent labeled ligands)
ΙT
     P-qlycoproteins
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (peptides and colchicine and vinblastine transport by; protein
        transport assays using IR fluorescent labeled ligands)
     Immunoglobulin receptors
IT
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (polymeric Ig, in detection of drugs affecting
        transport of IgA across epithelial cell membranes; protein transport
        assays using IR fluorescent labeled ligands)
     Biological transport
IΤ
     Drug screening
     Endocytosis
     Exocytosis
     Fluorescence
     Membrane, biological
     Microtiter plates
     Transcytosis
        (protein transport assays using IR fluorescent labeled ligands
        )
ΙT
     Receptors
     Transferrin receptors
     Transferrins
     Transport proteins
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (protein transport assays using IR fluorescent labeled ligands
ΙT
     Fluorometry
        (scanning; protein transport assays using IR fluorescent labeled
        ligands)
     Peptides, biological studies
TT
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (transport of, by P-glycoprotein; protein transport assays using IR
        fluorescent labeled ligands)
     29816-01-1D, conjugates with fluorescent labels
IT
     RL: ARG (Analytical reagent use); BPR (Biological process); BSU
     (Biological study, unclassified); ANST (Analytical study); BIOL
     (Biological study); PROC (Process); USES (Uses)
        (PEPT1 in pancreatic cells transport of; protein transport assays using
        IR fluorescent labeled ligands)
     166547-11-1D, NN382, conjugates with IgA
                                                166799-10-6D, IRD41, conjugates
ΙT
     with transferrin
                        169799-14-8D, Cy7, conjugates with transferrin
     172777-84-3D, Cy5.5, conjugates with IgA
     RL: ARG (Analytical reagent use); BPR (Biological process); BSU
     (Biological study, unclassified); ANST (Analytical study); BIOL
     (Biological study); PROC (Process); USES (Uses)
        (protein transport assays using IR fluorescent labeled ligands
        )
```

```
9004-54-0, Dextran, biological studies
ΤТ
    RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (protein transport assays using IR fluorescent labeled ligands
     52-53-9D, Verapamil, conjugates with fluorescent labels
                                                               64-86-8D.
IT
    Colchicine, conjugates with fluorescent labels
                                                      865-21-4D, Vinblastine,
                                        186042-32-0D, conjugates with
     conjugates with fluorescent labels
     fluorescent labels
     RL: ARG (Analytical reagent use); BPR (Biological process); BSU
     (Biological study, unclassified); ANST (Analytical study); BIOL
     (Biological study); PROC (Process); USES (Uses)
        (transport of, by P-glycoprotein; protein transport assays using IR
        fluorescent labeled ligands)
                                                865-21-4, Vinblastine
     52-53-9, Verapamil
                          64-86-8, Colchicine
TΤ
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (transport of, by P-glycoprotein; protein transport assays using IR
        fluorescent labeled ligands)
     51-41-2D, Noradrenaline, conjugates with fluorescent labels
ΙT
     RL: ARG (Analytical reagent use); BPR (Biological process); BSU
     (Biological study, unclassified); ANST (Analytical study); BIOL
     (Biological study); PROC (Process); USES (Uses)
        (transport of, through noradrenaline transporter; protein transport
        assays using IR fluorescent labeled ligands)
ΙT
     51-41-2, Noradrenaline
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (transport of, through noradrenaline transporter; protein transport
        assays using IR fluorescent labeled ligands)
              THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT
RE
(1) Cardonne; The Journal of Cell Biology 1994, V124(5), P717
(2) Lee; N-heteroaromatic ion and iminium ion substituted cyanine dyes for use
    as fluorescence labels
(3) Lipowska; Synthetic Communications 1993, V23(21), P3087 HCAPLUS
(4) Yue; US 5656449 A 1997 HCAPLUS
(5) Yue; US 5658751 A 1997 HCAPLUS
L66 ANSWER 5 OF 10 HCAPLUS COPYRIGHT 2003 ACS on STN
     2000:650656 HCAPLUS
ΑN
     134:206320
DN
     Role of J chain in secretory immunoglobulin formation
TI
ΑU
     Johansen, F.-E.; Braathen, R.; Brandtzaeg, P.
     Laboratory for Immunohistochemistry and Immunopathology (LIIPAT),
CS
     Institute of Pathology, University of Oslo, Rikshospitalet, Oslo, N-0027,
     Scandinavian Journal of Immunology (2000), 52(3), 240-248
SO
     CODEN: SJIMAX; ISSN: 0300-9475
     Blackwell Science Ltd.
PΒ
DT
     Journal
LA
     English
     15-3 (Immunochemistry)
CC
     The joining (J) chain is a small polypeptide, expressed by mucosal and
AB
     glandular plasma cells, which regulates polymer formation of IgA
     and IgM. J-chain incorporation into polymeric IgA (pIgA, mainly
     dimers) and pentameric IgM endows these antibodies with several salient
     features. First, a high valency of antigen-binding sites, which makes
     them suitable for agglutinating bacteria and viruses; little or no
     complement-activating potential, which allows them to operate in a
     noninflammatory fashion; and, most importantly, only J-chain-contg.
     polymers show high affinity for the polymeric Ig
```

receptor (pIgR), also known as transmembrane

STΙT

ΙT

IT

ΙT

ΙT

ΙT

IT

IT

ΙT

IT

IT

secretory component (SC). This epithelial glycoprotein mediates active external transfer of pIgA and pentameric IgM to exocrine secretions. Thus, secretory IgA (SIgA) and SIgM, as well as free SC, are generated by endoproteolytic cleavage of the pIgR extracellular domain. The secretory antibodies form the "first line" of defense against pathogens and noxious substances that favor the mucosae as their portal of entry. The J chain is involved in creating the binding site for plgR/SC in the lg polymers, not only by detg. the polymeric quaternary structure but apparently also by interacting directly with the receptor protein. Therefore, both the J chain and the plgR/SC are key proteins in secretory immunity. J chain secretory IgA IgM Immunoglobulins RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (A, polymeric; J chain role in secretory Ig formation) Immunoglobulins RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (A, secretory; J chain role in secretory Ig 'formation) -B cell (lymphocyte) (J chain role in secretory Ig formation in) Immunoglobulins RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (M, pentameric; J chain role in secretory Ig formation) Immunoglobulins RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (M, secretory; J chain role in secretory Ig formation) Immunoglobulins RL: BAC (Biological activity or effector, except adverse); BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (fragments; J chain role in secretory Ig formation) Lymphocyte (plasma cell; J chain role in secretory Ig formation in) Immunoglobulin receptors RL: BOC (Biological occurrence); BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); OCCU (Occurrence); PROC (Process) (polymeric Ig; J chain role in secretory Ig formation) Quaternary structure (protein; J chain role in secretory Ig formation) Immunoglobulins RL: BOC (Biological occurrence); BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); OCCU (Occurrence); PROC (Process) (secretory component; J chain role in secretory Ig formation) Immunity (secretory; J chain role in secretory Ig formation in relation to) THERE ARE 85 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 85 (1) Atkin, J; J Immunol 1996, V157, P156 HCAPLUS (2) Bakos, M; J Immunol 1993, V151, P1346 HCAPLUS (3) Bastian, A; Adv Exp Med Biol 1995, V371A, P581 HCAPLUS (4) Boehm, M; J Mol Biol 1999, V286, P1421 HCAPLUS

Page 13

```
(5) Brandtzaeg, P; Adv Exp Med Biol 1974, V45, P87 HCAPLUS
(6) Brandtzaeg, P; Ann N Y Acad Sci 1996, V778, P1 HCAPLUS
(7) Brandtzaeg, P; Clin Exp Immunol 1984, V58, P709 HCAPLUS
(8) Brandtzaeg, P; Immunochemistry 1977, V14, P179 HCAPLUS
(9) Brandtzaeg, P; Immunol Rev 1999, V171, P45 HCAPLUS
(10) Brandtzaeg, P; Immunol Today 1999, V20, P141 HCAPLUS
(11) Brandtzaeg, P; Immunology 1975, V29, P559 HCAPLUS
(12) Brandtzaeg, P; Mol Immunol 1983, V20, P941 HCAPLUS
(13) Brandtzaeg, P; Nature 1968, V220, P292 HCAPLUS
(14) Brandtzaeg, P; Nature 1974, V252, P418 MEDLINE
(15) Brandtzaeg, P; Nature 1984, V311, P71 HCAPLUS
(16) Brandtzaeg, P; Nature New Biol 1973, V243, P142 HCAPLUS
(17) Brandtzaeg, P; Scand J Immunol 1975, V4, P439 HCAPLUS
(18) Brandtzaeg, P; Scand J Immunol 1975, V4, P837 HCAPLUS
(19) Brandtzaeg, P; Scand J Immunol 1976, V5, P411 HCAPLUS
(20) Brandtzaeg, P; Scand J Immunol 1985, V22, P111 HCAPLUS
(21) Brewer, J; Mol Immunol 1997, V34, P323 HCAPLUS
(22) Cattaneo, A; EMBO J 1987, V6, P2753 HCAPLUS
(23) Coyne, R; J Biol Chem 1994, V269, P31620 HCAPLUS
(24) Crottet, P; Biochem J 1999, V341, P299 HCAPLUS
(25) Davis, A; EMBO J 1989, V8, P2519 HCAPLUS
(26) Davis, A; Eur J Immunol 1988, V18, P1001 HCAPLUS
(27) Davis, A; J Immunol 1989, V143, P1352 HCAPLUS
(28) Della Corte, E; Biochem J 1973, V136, P597 HCAPLUS
(29) Dickinson, B; J Clin Invest 1999, V104, P903 HCAPLUS
(30) Erlandsson, L; Eur J Immunol 1998, V28, P2355 HCAPLUS
(31) Eskeland, T; Immunochemistry 1974, V11, P161 HCAPLUS
(32) Fallgreen-Gebauer, E; Biol Chem Hoppe-Seyler 1993, V374, P1023 HCAPLUS
(33) Frutiger, S; Biochemistry 1992, V31, P12643 HCAPLUS
(34) Garcia-Pardo, A; J Biol Chem 1981, V256, P11734 HCAPLUS
(35) Geneste, C; Immunol Let 1986, V13, P221 HCAPLUS
(36) Grubb, A; Acta Med Scand 1978, V204, P453 HCAPLUS
(37) Halpern, M; Nature 1970, V228, P1276 HCAPLUS
(38) Hendrickson, B; J Exp Med 1995, V182, P1905 HCAPLUS
(39) Hendrickson, B; J Immunol 1996, V157, P750 HCAPLUS
(40) Hexham, J; J Exp Med 1999, V189, P747 HCAPLUS
(41) Hohman, V; Mol Immunol 1997, V34, P995 HCAPLUS
(42) Hughes, G; Biochem J 1990, V271, P641 HCAPLUS
(43) Johansen, F; Eur J Immunol 1999, V29, P1701 HCAPLUS
(44) Johansen, F; J Exp Med 1999, V190, P915 HCAPLUS
(45) Koshland, M; J Immunol 1977, V118, P775 HCAPLUS
(46) Krugmann, S; J Immunol 1997, V159, P244 HCAPLUS
(47) Kulseth, M; DNA Cell Biol 1994, V13, P37 HCAPLUS
(48) Lycke, N; J Immunol 1999, V163, P913 HCAPLUS
(49) Matsuuchi, L; Proc Natl Acad Sci USA 1986, V83, P456 HCAPLUS
(50) Max, E; J Exp Med 1985, V161, P832 HCAPLUS
(51) Max, E; Proc Natl Acad Sci USA 1986, V83, P5592 HCAPLUS
(52) Mestecky, J; Nature 1974, V249, P650 HCAPLUS
(53) Mestecky, J; Proc Natl Acad Sci USA 1974, V71, P544 HCAPLUS
(54) Mestecky, J; Science 1971, V171, P1163 HCAPLUS
(55) Metzger, H; Adv Immunol 1970, V12, P57 HCAPLUS
(56) Mosmann, T; Eur J Immunol 1978, V8, P94 HCAPLUS
(57) Mostov, K; Mucosal Immunology 1999, P181
(58) Niles, M; Proc Natl Acad Sci USA 1995, V92, P2884 HCAPLUS
(59) Norderhaug, I; Crit Rev Immunol 1999, V19, P481 HCAPLUS
(60) Norderhaug, I; Eur J Immunol 1999, V29, P3401 HCAPLUS
(61) Parkhouse, R; Immunology 1970, V18, P575 HCAPLUS
(62) Radl, J; Immunology 1971, V20, P843 HCAPLUS
(63) Randall, T; Eur J Immunol 1990, V20, P1971 HCAPLUS
(64) Randall, T; J Biol Chem 1992, V267, P18002 HCAPLUS
(65) Randall, T; Proc Natl Acad Sci USA 1992, V89, P962 HCAPLUS
```

(66) Roe, M; J Immunol 1999, V162, P6046 HCAPLUS (67) Roth, R; Biochemistry 1981, V20, P6594 HCAPLUS

```
(68) Russell, M; Eur J Immunol 1989, V19, P2243 HCAPLUS
(69) Shimada, S; J Immunol 1999, V163, P5367 HCAPLUS
(70) Sitia, R; Cell 1990, V60, P781 HCAPLUS
(71) Socken, D; Immunochemistry 1978, V15, P499 HCAPLUS
(72) Sorensen, V; Int Immunol 2000, V12, P19 HCAPLUS
(73) Sorensen, V; J Immunol 1996, V156, P2858 MEDLINE (74) Sorensen, V; J Immunol 1999, V162, P3448 HCAPLUS
(75) Takahashi, T; Immunogenetics 2000, V51, P85 HCAPLUS
(76) Takahashi, T; Proc Natl Acad Sci USA 1996, V93, P1886 HCAPLUS
(77) Tomasi, T; J Exp Med 1965, V121, P101 HCAPLUS
(78) Underdown, B; Ann Rev Immunol 1986, V4, P389 HCAPLUS
(79) Vaerman, J; Eur J Immunol 1998, V28, P171 HCAPLUS
(80) Vaerman, J; Immunol Invest 1995, V24, P631 HCAPLUS
(81) Vaerman, J; Immunology 1998, V95, P90 HCAPLUS
(82) Wiersma, E; J Immunol 1998, V160, P5979 HCAPLUS
(83) Yoo, E; J Biol Chem 1999, V274, P33771 HCAPLUS
(84) Zikan, J; Mol Immunol 1986, V23, P541 HCAPLUS
(85) Zikan, J; Proc Natl Acad Sci USA 1985, V82, P5905 HCAPLUS
     ANSWER 6 OF 10 HCAPLUS COPYRIGHT 2003 ACS on STN
L66
     2000:220716 HCAPLUS
AN
DN
     132:261375
     Immunoglobulin fusion product with immunoglobulin receptor that protects
ΤI
     Ig in mucosal environment, cDNA sequences, transgenic plants, and dental
     caries prevention
     Hiatt, Andrew C.; Ma, Julian K. C.; Lehner, Thomas; Mostov, Keith
ΙN
PA
     USA
     U.S., 59 pp., Cont.-in-part of U.S. Ser. No. 367,395.
SO
     CODEN: USXXAM
DT
     Patent
LA
     English
IC
     ICM C12N015-00
         C12N015-29; C12N015-82; A01H004-00
NCL
     435070100
     3-2 (Biochemical Genetics)
     Section cross-reference(s): 1, 11, 15
FAN.CNT 2
                      KIND
                            DATE
                                            APPLICATION NO.
                                                              DATE
     PATENT NO.
     -----
                             _____
                      ____
     US 6046037
                             20000404
                                            US 1995-434000
                                                              19950504
PΙ
                       Α
                       AΑ
                             19960711
                                            CA 1995-2208783
                                                              19951227
     CA 2208783
                                            WO 1995-US16889
                                                              19951227
     WO 9621012
                       Α1
                             19960711
         W: AU, BR, CA, CN, CZ, FI, HU, JP, KR, MX, NO, NZ, PL, RU, SG
         RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
                             19960724
                                            AU 1996-46088
                                                              19951227
     AU 9646088
                       Α1
     AU 722668
                       B2
                             20000810
                             19971119
                                            EP 1995-944237
                                                              19951227
     EP 807173
                       A1
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE
                             19980603
                                            CN 1995-197699
                                                              19951227
     CN 1183802
                       Α
                                            US 1999-312157
                                                              19990514
     US 6303341
                       В1
                             20011016
                                            US 2001-982107
                                                              20011016
     US 2002159958
                       A1
                             20021031
PRAI US 1994-367395
                       B2
                             19941230
     US 1995-434000
                       Α
                             19950504
                       W
                             19951227
     WO 1995-US16889
                       A1
                             19990514
     US 1999-312157
     Igs of the present invention are useful as therapeutic Igs against mucosal
AB
     pathogens such as Streptococcus mutans. The Igs contain a protection
     protein (e.g., the polyimmunoglobulin receptor) that protects the Igs in
     the mucosal environment. The invention also includes a greatly improved
     method of producing Igs in plants by producing the protection protein in
```

the same cell as the other components of the Igs. The components of the

Ig are assembled at a much improved efficiency. The method of the

invention allows the assembly and high efficiency prodn. of such complex mols. The invention also contemplates the prodn. of Igs contg. protection proteins in a variety of cells, including plant cells, that can be selected for useful addnl. properties. The use of Igs contg. protection proteins as therapeutic **antibodies** against mucosal and other pathogens is also contemplated.

- ST Ig fusion receptor protection mucosa caries; dental caries prevention Ig fusion receptor; sequence Ig fusion receptor mucosa protection; plant transgenic manuf Ig fusion receptor
- IT Immunoglobulins
 - RL: BMF (Bioindustrial manufacture); BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
 - (A; Ig fusion product with Ig receptor that protects Ig in mucosal environment, cDNA sequences, transgenic plants, and dental caries prevention)
- IT Immunoglobulins
 - RL: BMF (Bioindustrial manufacture); BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
 - (D; Ig fusion product with Ig receptor that protects Ig in mucosal environment, cDNA sequences, transgenic plants, and dental caries prevention)
- IT Immunoglobulins
 - RL: BMF (Bioindustrial manufacture); BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
 - (E; Ig fusion product with Ig receptor that protects Ig in mucosal environment, cDNA sequences, transgenic plants, and dental caries prevention)
- IT Immunoglobulins
 - RL: BMF (Bioindustrial manufacture); BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
 - (G; Ig fusion product with Ig receptor that protects Ig in mucosal environment, cDNA sequences, transgenic plants, and dental caries prevention)
- IT Immunoglobulins
 - RL: BMF (Bioindustrial manufacture); BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
 - (Guy's 13, fusion products, with Ig receptors; Ig fusion product with Ig receptor that protects Ig in mucosal environment, cDNA sequences, transgenic plants, and dental caries prevention)
- IT Antigens
 - RL: BSU (Biological study, unclassified); BIOL (Biological study)
 (Ig antigen-binding domain; Ig fusion product with Ig receptor that
 protects Ig in mucosal environment, cDNA sequences, transgenic plants,
 and dental caries prevention)
- IT Agrobacterium tumefaciens
 - Alfalfa (Medicago sativa)
 - Arabidopsis
 - Dicotyledon (Magnoliopsida)
 - Immunotherapy
 - Monocotyledon (Liliopsida)
 - Mucous membrane
 - Petunia
 - Protein sequences
 - Streptococcus mutans
 - Streptococcus sobrinus
 - Tobacco
 - Tomato
 - cDNA sequences

(Ig fusion product with Ig receptor that protects Ig in mucosal environment, cDNA sequences, transgenic plants, and dental caries prevention)

IT Immunoglobulins

RL: BMF (Bioindustrial manufacture); BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(M; Ig fusion product with Ig receptor that protects Ig in mucosal environment, cDNA sequences, transgenic plants, and dental caries prevention)

IT Tooth

(caries, prevention of; Ig fusion product with Ig receptor that protects Ig in mucosal environment, cDNA sequences, transgenic plants, and dental caries prevention)

IT Immunoglobulin receptors

RL: BMF (Bioindustrial manufacture); BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(fusion products, with Igs; Ig fusion product with Ig receptor that protects Ig in mucosal environment, cDNA sequences, transgenic plants, and dental caries prevention)

IT Transformation, genetic

(transgenic, Ig fusion product with Ig receptor that protects Ig in mucosal environment, cDNA sequences, transgenic plants, and dental caries prevention)

- 144997-23-9DP, Glycoprotein (human secretory component protein TT moiety reduced), fusion products with Ig 170979-93-8DP, fusion products 180616-69-7DP, Receptor, immunoglobulin (rabbit), fusion 180616-70-0DP, Receptor, immunoglobulin (mouse), fusion products with Iq 180686-83-3DP, Receptor, immunoglobulin (rat), fusion products with Iq 180686-85-5DP, fusion products with Ig receptor products with Ig 180686-87-7DP, fusion products with Ig receptor RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (amino acid sequence; Ig fusion product with Ig receptor that protects Ig in mucosal environment, cDNA sequences, transgenic plants, and dental caries prevention)
- IT 140080-71-3DP, fusion products with Ig 140262-61-9DP, fusion products with Ig 153420-82-7DP, fusion products with Ig 153665-28-2DP, fusion products with Ig 159070-18-5DP, fusion products with Ig 180686-84-4DP, fusion products with Ig receptor cDNA 180686-86-6DP, fusion products with Ig receptor cDNA

RL: BPN (Biosynthetic preparation); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (nucleotide sequence; Ig fusion product with Ig receptor that protects Ig in mucosal environment, cDNA sequences, transgenic plants, and dental caries prevention)

IT 244135-92-0, PN: US5959177 SEQID: 13 unclaimed DNA 244135-94-2, PN: US5959177 SEQID: 14 unclaimed DNA 244135-95-3, PN: US5959177 SEQID: 15 unclaimed DNA 244135-96-4, PN: US5959177 SEQID: 16 unclaimed DNA 244135-97-5, PN: US5959177 SEQID: 17 unclaimed DNA RL: PRP (Properties)

(unclaimed nucleotide sequence; Ig fusion product with Ig receptor that protects Ig in mucosal environment, cDNA sequences, transgenic plants, and dental caries prevention)

RE.CNT 95 THERE ARE 95 CITED REFERENCES AVAILABLE FOR THIS RECORD RE

- (1) Abdullah, R; Biotechnology 1986, V4, P1087
- (2) Anon; WO 8700551 1987 HCAPLUS
- (3) Anon; WO 9014430 1990 HCAPLUS
- (4) Anon; EP 480014 B1 1991 HCAPLUS
- (5) Anon; WO 9106320 1991 HCAPLUS
- (6) Anon; WO 9116061 1991 HCAPLUS

- (7) Anon; EP 484148 A1 1992
- (8) Anon; EP 0371017 B1 1994 HCAPLUS
- (9) Anon; Inaugural Dissertation 1988
- (10) Anon; Plant Molecular Biology 1990, V15, P281
- (11) Anon; Plant Molecular Biology 1990, V15, P281
- (12) Bakos; Molecular Immunology 1994, V31(2), P165 HCAPLUS
- (13) Banting, G; FEBS Letters 1989, V254, P177 HCAPLUS
- (14) Barnes, W; Proc Natl Acad Sci USA 1990, V87, P9183 HCAPLUS
- (15) Benbrook, C; Proceedings Bio Expo 1986, P27
- (16) Benfey, P; Science 1989, V244, P174 HCAPLUS
- (17) Benfey, P; Science 1990, V250, P959 HCAPLUS
- (18) Brandtzaeg, P; Nature 1984, V311, P71 HCAPLUS
- (19) Bytebier, B; Proc Natl Acad Sci USA 1987, V84, P5345 HCAPLUS
- (20) Callis, J; Genes and Development 1987, V1, P1183 HCAPLUS
- (21) Carayannopoulos, L; Proc Natl Acad Sci USA 1994, V91, P8348 HCAPLUS
- (22) Cocking; Science 1987, V236, P1259
- (23) Corthesy, B; Experientai 1994, V50, PA27
- (24) Crago; Journal of Immunology 1989, V142(11), P3909 HCAPLUS
- (25) de la Pena, A; Nature 1987, V325, P274 HCAPLUS
- (26) Eliasson; Journal of Biological Chemistry 1989, V263(9), P4323
- (27) Evans; US 4870009 1989 HCAPLUS
- (28) Fischhoff; US 5349124 1994 HCAPLUS
- (29) Fraley; US 5352605 1994 HCAPLUS
- (30) Fraley, R; Proc Natl Acad Sci USA 1983, V80, P4803 HCAPLUS
- (31) Fromm, M; Nature 1986, V319, P791 HCAPLUS
- (32) Gilchrest; US 5352440 1994 HCAPLUS
- (33) Hein, M; Biotechnol Prog 1991, V7, P455 HCAPLUS
- (34) Hess, D; International Review of Cytology 1987, V107, P367
- (35) Hiatt; Intern Rev Immunol 1993, V10, P139 MEDLINE
- (36) Hiatt, A; FEBS Letters 1992, V307(1), P71 HCAPLUS
- (37) Hiatt, A; Nature 1989, V342, P76 HCAPLUS
- (38) Hiatt, A; The Pharmacology of Monoclonal Antibodies 1994, Chapter 12, P317
- (39) Horsch, R; Science 1985, V227, P1229 HCAPLUS
- (40) Huang, A; Cell 1981, V27, P245 HCAPLUS
- (41) Huse, W; Science 1989, V246, P1275 HCAPLUS
- (42) Jorgensen, R; Mol Gen Genet 1987, V207, P471 HCAPLUS
- (43) Klein, T; Nature 1987, V327, P70 HCAPLUS
- (44) Klein, T; Proc Natl Acad Sci USA 1988, V85, P8502 HCAPLUS
- (45) Kobayashi, K; Immunochemistry 1973, V10, P73 HCAPLUS
- (46) Koiyumaki; US 4607388 1986
- (47) Koshland, M; Immunoglobulin Genes 1989, Chap 18, P345
- (48) Kraehenbuhl, J; Trends in Cell Biol 1992, V2, P170 HCAPLUS
- (49) Kraehenbul; Advances in Experimental Medicine and Biology 1987, V216B, P1053
- (50) Krajci, P; Biochem Biophys Res Comm 1989, V158, P783 HCAPLUS
- (51) Krajci, P; Eur J Immunol 1992, V22, P2309 HCAPLUS
- (52) Lambda; Science 1989, V246, P1275
- (53) Larrick, J; The Pharmacology of Monoclonal Antibodies 1994, Chapter 2, P23
- (54) Leder; US 4736866 1988 HCAPLUS
- (55) Lee, C; Infection and Immunity 1994, V62(3), P887 HCAPLUS
- (56) Lehner; US 4594244 1986 HCAPLUS
- (57) Lehner; US 5352446 1994 HCAPLUS
- (58) Lindh, E; The Journal of Immunology 1975, V114(1), P284
- (59) Lorz, H; Mol Gen Genet 1985, V199, P178
- (60) Luo, Z; Plant Mol Biol Reporter 1988, V6, P165 HCAPLUS
- (61) Ma, J; Clin Exp Immunol 1989, V77, P331 MEDLINE
- (62) Ma, J; Eur J Immunol 1994, V24, P131 HCAPLUS
- (63) Ma, J; Science 1995, V268, P716 HCAPLUS
- (64) Marcotte, W; Nature 1988, V335, P454 HCAPLUS
- (65) Mark, G; The Pharmacology of Monoclonal Antibodies 1994, Chapter 4, P105
- (66) Marshall, R; Annual Review of Biochemistry 1972, V41, P673 HCAPLUS
- (67) Marshall, R; Biochem Soc Symp 1974, V40, P17 HCAPLUS
- (68) Matsuuchi, L; Proc Natl Acad Sci USA 1986, V83, P456 HCAPLUS

```
(69) McCabe, D; Biotechnology 1988, V6, P923
(70) McNabb, P; Ann Rev Microbiol 1981, V35, P477 HCAPLUS
(71) Mostov, K; Ann Rev Immol 1994, V12, P63 HCAPLUS
(72) Mostov, K; Nature 1984, V308, P37 HCAPLUS
(73) Neuhaus, G; Theor Appl Genet 1987, V75, P30
(74) Odell, J; Nature 1985, V313, P810 HCAPLUS
(75) Orlandi, R; Proc Natl Acad Sci USA 1989, V86, P3833 HCAPLUS
(76) Paszkowski, J; The EMBO Journal 1989, V3, P2717
(77) Piskurich, J; J Immunol 1993, V150, P38
(78) Potrykus, I; Mol Gen Genet 1985, V199, P183 HCAPLUS
(79) Rogers; US 5034322 1991 HCAPLUS
(80) Rogers, S; Methods in Enzymology 1987, V153, P253 HCAPLUS
(81) Sadowski; US 4443549 1984 HCAPLUS
(82) Sadowski; US 4652448 1987 HCAPLUS
(83) Schlom; US 5183756 1993 HCAPLUS
(84) Shah; US 5188642 1993 HCAPLUS
(85) Silbart, L; Science 1989, V243, P1462 HCAPLUS
(86) Smith, R; Oral Microbiol Immunol 1989, V4, P153 HCAPLUS
(87) Solari; Biochemical Journal 1989, V257, P759 HCAPLUS
(88) Spielmann, A; Mol Gen Genet 1986, V205, P34 HCAPLUS
(89) Toriyama, K; Theor Appl Genet 1986, V73, P16
(90) Uchimiya, H; Mol Gen Genet 1986, V204, P204 HCAPLUS
(91) Vasil, I; Biotechnology 1988, V6, P397
(92) Verbeet; GenBank Accession No X81371
(93) Wagner; US 4873191 1989
(94) Williams, A; Immunoglobulin Genes 1989, Chap 19, P361
(95) Zhou, G; Methods in Enzymology 1983, V101, P433 HCAPLUS
L66
     ANSWER 7 OF 10 HCAPLUS COPYRIGHT 2003 ACS on STN
     1998:1507 HCAPLUS
AN
DN
     128:74313
     Cellular internalization of the polymeric Iq
ΤI
     receptor and of antibody ligands directed to
     the extracellular pIgR stalk
    Mostov, Keith E.; Richman-Eisenstat, Janice
ΙN
     Regents of the University of California, USA
PΑ
     PCT Int. Appl., 41 pp.
SO
     CODEN: PIXXD2
DT
     Patent
LA
     English
     ICM C07K016-00
IC
     15-3 (Immunochemistry)
     Section cross-reference(s): 1
FAN.CNT 1
     PATENT NO.
                      KIND DATE
                                           APPLICATION NO.
                                                            DATE
                                                           _____
                                           -----
     ------
                      ----
                           -----
                                           WO 1997-US7944 19970514
                     A1 19971211
     WO 9746588
ΡI
            AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE,
             DK, EE, ES, FI, GB, GE, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LC,
             LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT,
             RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, UZ, VN, YU,
             AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FI, FR, GB,
             GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN,
             ML, MR, NE, SN, TD, TG
                                                            19970514
                                           CA 1997-2256304
     CA 2256304
                       AA
                            19971211
                                                            19970514
     AU 9730632
                            19980105
                                           AU 1997-30632
                       A1
     AU 728587
                       B2
                            20010111
                                           CN 1997-195238
                                                            19970514
     CN 1221428
                       Α
                            19990630
     EP 934338
                       A1
                            19990811
                                           EP 1997-925515
                                                            19970514
             AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, FI
                            20000328
                                           US 1997-856383
                                                            19970514
     US 6042833
```

```
JP 1998-500584
                            20000905
                                                            19970514
    JP 2000511432
                       Т2
    IL 127238
                      A1
                                                            19970514
                            20010724
                                           IL 1997-127238
                      C2
                            20021027
    RU 2191781
                                           RU 1999-100279
                                                            19970514
                      В1
                            20020122
                                           US 1999-475088
                                                            19991230
    US 6340743
                      P
PRAI US 1996-18958P
                            19960604
                      A3
    US 1997-856383
                            19970514
                      W
                            19970514
    WO 1997-US7944
    The present invention is directed to a liqand that binds
AΒ
    specifically to the stalk of a polymeric Ig
    receptor (pIgR) of a cell in a secretory
    component-independent manner. Disclosed are methods of attaching and
    introducing a ligand into a cell expressing pIgR. The
    invention provides the means for transporting therapeutic or diagnostic
    compns. to, into (endocytosis) or across a cell expressing plqR.
ST
    internalization polymeric Ig receptor
    ligand
    Gene, animal
TΤ
    RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (CFTR; cellular internalization of polymeric Iq
        receptor and of antibody ligands joined to)
    Immunoglobulins
ΙT
    RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (Y; cellular internalization of polymeric Ig
        receptor and of antibody ligands directed
        to extracellular pIgR stalk)
ΙT
    Diagnosis
    Gene therapy
    Immunotherapy
    Transcytosis
        (cellular internalization of polymeric Ig
        receptor and of antibody ligands directed
        to extracellular pIgR stalk)
ΙT
    Antibodies
    RL: BPR (Biological process); BSU (Biological study, unclassified); THU
     (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses)
        (cellular internalization of polymeric Ig
        receptor and of antibody ligands directed
        to extracellular pIgR stalk)
IT
    Anti-infective agents
    Anti-inflammatory agents
    Antibiotics
    Plasmids
        (cellular internalization of polymeric Ig
        receptor and of antibody ligands joined to)
ΙT
    Antisense oligonucleotides
    Carbohydrates, biological studies
    Lipids, biological studies
    Nucleic acids
    Proteins, specific or class
    Radionuclides, biological studies
    RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (cellular internalization of polymeric Ig
        receptor and of antibody ligands joined to)
    CFTR (cystic fibrosis transmembrane conductance regulator)
IT
    RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (cellular internalization of polymeric Ig
        receptor and of antibody ligands joined to
        gene for)
IT
     Digestive tract
     Respiratory tract
        (epithelium; cellular internalization of polymeric Ig
        receptor and of antibody ligands directed
```

```
to extracellular pIgR stalk)
     Immunoglobulins
IT
     RL: BPR (Biological process); BSU (Biological study, unclassified); THU
     (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses)
        (fragments, conjugates, with polylysine; cellular
        internalization of polymeric Ig receptor
        and of antibody ligands directed to extracellular
        pIgR stalk)
     Antibodies
TΤ
     RL: BPR (Biological process); BSU (Biological study, unclassified); THU
     (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses)
        (humanized; cellular internalization of polymeric Ig
        receptor and of antibody ligands directed
        to extracellular pIgR stalk)
TT
     Animal cell
        (mammalian; cellular internalization of polymeric Ig
        receptor and of antibody ligands directed
        to extracellular pIgR stalk)
IT
     Protein sequences
        (of polymeric Ig receptors of mammals)
IT
     Immunoglobulin receptors
     RL: BPR (Biological process); BSU (Biological study, unclassified); PRP
     (Properties); THU (Therapeutic use); BIOL (Biological study); PROC
     (Process); USES (Uses)
        (polymeric Ig; cellular internalization of
        polymeric Ig receptor and of
        antibody ligands directed to extracellular
        pIgR stalk)
IT
     Endocytosis
        (receptor-mediated; cellular internalization of
        polymeric Ig receptor and of
        antibody ligands directed to extracellular
        pIgR stalk)
     Biological transport
TT
        (retrograde; cellular internalization of polymeric Ig
        receptor and of antibody ligands directed
        to extracellular pIgR stalk)
ΤТ
     Antibodies
     RL: BPR (Biological process); BSU (Biological study, unclassified); THU
     (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses)
        (single chain; cellular internalization of polymeric
        Ig receptor and of antibody ligands
        directed to extracellular plgR stalk)
     25104-18-1D, Poly-L-lysine, antibody Fab conjugates
ΙT
     RL: BPR (Biological process); BSU (Biological study, unclassified); THU
     (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses)
        (cellular internalization of polymeric Ig
        receptor and of antibody ligands directed
        to extracellular pIgR stalk)
                                               200392-09-2
                                                              200513-53-7
                                 200392-08-1
ΙT
     200392-06-9
                   200392-07-0
     200578-06-9
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (cellular internalization of polymeric Ig
        receptor and of antibody ligands targeted
        to)
L66 ANSWER 8 OF 10 HCAPLUS COPYRIGHT 2003 ACS on STN
AN
     1996:482531 HCAPLUS
     125:191224
DN
     Regulation of protein traffic in polarized epithelial cells: the
TΤ
     polymeric immunoglobulin receptor model
     Mostov, K. E.; Altschuler, Y.; Chapin, S. J.; Enrich,
ΑU
```

```
C.; Low, S.-H.; Luton, F.; Richman-Eisenstat, J.; Singer, K. L.;
     Tang, K.; Weimbs, T.
     Department Anatomy, University California, San Francisco, CA, 94143-0452,
CS
     USA
     Cold Spring Harbor Symposia on Quantitative Biology (1995), 60(Protein
SO
     Kinesis: The Dynamics of Protein Trafficking and Stability), 775-781
     CODEN: CSHSAZ; ISSN: 0091-7451
     Cold Spring Harbor Laboratory Press
PΒ
     Journal; General Review
DT
LA
     English
CC
     13-0 (Mammalian Biochemistry)
     A review with 25 refs. The polymeric Ig
ΑB
     receptor (pIgR) provides an excellent model for
     analyzing the regulation of membrane traffic in polarized epithelial
     cells. The basolateral sorting signal of the pIgR and the
     pathway and regulation of transcytosis were discussed in detail.
     review protein transport polarized epithelium; polymeric
ST
     Iq receptor transport epithelium review
     Proteins, biological studies
ΙT
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (polymeric Ig receptor model of
        regulation of protein traffic in polarized epithelial cells)
IT
     Immunoglobulin receptors
      Receptors
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (pIgR (polymeric Ig receptors),
        polymeric Ig receptor model of regulation
        of protein traffic in polarized epithelial cells)
ΙT
     Epithelium
        (polarized, polymeric Ig receptor model
        of regulation of protein traffic in polarized epithelial cells)
ΙT
     Biological transport
        (translocation, polymeric Ig receptor
        model of regulation of protein traffic in polarized epithelial cells)
    ANSWER 9 OF 10 HCAPLUS COPYRIGHT 2003 ACS on STN
AN
     1996:54805 HCAPLUS
DN
     124:115363
     Calmodulin binds to the basolateral targeting signal of the
TI
     polymeric immunoglobulin receptor
     Chapin, Steven J.; Enrich, Carlos; Aroeti, Benjamin; Havel,
ΑU
     Richard J.; Mostov, Keith E.
     Dep. Anat. Biochem. Biophys., Univ. California, San Francisco, CA, 94143,
CS
     USA
     Journal of Biological Chemistry (1996), 271(3), 1336-42
SO
     CODEN: JBCHA3; ISSN: 0021-9258
PΒ
     American Society for Biochemistry and Molecular Biology
DT
     Journal
LA
     English
CC
     15-10 (Immunochemistry)
     Section cross-reference(s): 13
     We have identified a major calmodulin (CaM)-binding protein in rat liver
AB
     endosomes using 125I-CaM overlays from two-dimensional protein blots.
     Immunostaining of blots demonstrates that this protein is the
     polymeric Ig receptor (pIgR). We
     further investigated the interaction between plgR and CaM using
     Madin-Darby canine kidney cells stably expressing cloned wild-type and
     mutant plqR. We found that detergent-solubilized plqR
     binds to CaM-agarose in a Ca2+-dependent fashion, and binding is inhibited
     by the addn. of excess free CaM or the CaM antagonist W-13 \,
     (N-(4-aminobutyl)-5-chloro-2-naphthalenesulfonamide), suggesting that
```

pIgR binding to CaM is specific. Furthermore, pIgR is the most prominent 35S-labeled CaM-binding protein in the detergent phase of Triton X-114-solubilized, metabolically labeled pIgR -expressing Madin-Darby canine kidney cells. CaM can be chem. cross-linked to both solubilized and membrane-assocd. pigR, suggesting that binding can occur while the pIgR is in intact membranes. The CaM binding site is located in the membrane-proximal 17-amino acid segment of the pIgR cytoplasmic tail. This region of pIgR constitutes an autonomous basolateral targeting signal. However, binding of CaM to various pIgR mutants suggests that CaM binding is not necessary for basolateral targeting. We suggest that CaM may be involved in regulation of plgR transcytosis and/or signaling by pIgR. calmodulin binding polymeric Ig receptor; binding site polymeric Ig receptor calmodulin Calmodulins RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (calmodulin binds to the basolateral targeting signal of the polymeric Ig receptor) Immunoglobulin receptors Receptors RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (pIgR (polymeric Ig receptors), calmodulin binds to the basolateral targeting signal of the polymeric Ig receptor) 7440-70-2, Calcium, biological studies RL: BSU (Biological study, unclassified); BIOL (Biological study) (calcium-dependent binding of calmodulin to the basolateral targeting signal of the polymeric Ig receptor) ANSWER 10 OF 10 HCAPLUS COPYRIGHT 2003 ACS on STN 1994:75120 HCAPLUS 120:75120 Stimulation of transcytosis of the polymeric immunoglobulin receptor by dimeric IgA Song, Wenxia; Bomsel, Morgane; Casanova, James; Vaerman, Jean Pierre; Mostov, Keith Cardiovasc. Res. Inst., Univ. California, San Francisco, CA, 94143-0452, USA Proceedings of the National Academy of Sciences of the United States of America (**1994**), 91(1), 163-6 CODEN: PNASA6; ISSN: 0027-8424 Journal English 15-3 (Immunochemistry) Section cross-reference(s): 6 The polymeric Ig receptor (pIgR) is transcytosed from the basolateral to the apical surface of polarized epithelial cells. The authors have previously shown that phosphorylation of Ser-664 in the cytoplasmic domain of the pIgR is a signal for its transcytosis. The authors now report that binding of a physiol. ligand, dimeric IgA, to pIgR stimulates pIgR transcytosis. This stimulation occurs in both the presence or absence of Ser-664 phosphorylation. The authors have used three methods to measure transcytosis of the plgR. The plgR was biosynthetically labeled and its cleavage to secretory component after transcytosis was measured. The pIgR was labeled with biotin at the basolateral surface. After transcytosis, release of the biotin-labeled secretory component into the apical medium was

measured. Transcytosis of a ligand bound to the pIgR

ST

IT

ΙT

IT

ΑN

DN

ΤI

ΑU

CS

SO

DT

LA

CC

AB

was measured. All three methods indicated that dimeric IgA stimulates transcytosis of the pIgR. dimeric IgA transcytosis polymeric Ig receptor STΙT Immunoglobulins RL: BIOL (Biological study) (A, dimers, polymeric Ig receptor transcytosis by polarized epithelium stimulation by) ΙT Kidney, metabolism (epithelium, polymeric Ig receptor transcytosis by, dimeric IgA ligand stimulation of) ΙT Receptors RL: BIOL (Biological study) (pIgR (polymeric Ig receptors), transcytosis of, by polarized epithelium, dimeric IgA ligand stimulation of) ΙT Immunoglobulins RL: BIOL (Biological study) (pigR receptors, transcytosis of, by polarized epithelium, dimeric IqA **ligand** stimulation of) ΙT Epithelium (polarized, polymeric Ig receptor transcytosis by, dimeric IgA ligand stimulation of) ΙT Biological transport (transcytosis, of polymeric Ig receptor, by polarized epithelium, dimeric IgA ligand stimulation of) => fil biosis FILE 'BIOSIS' ENTERED AT 07:20:20 ON 23 JUL 2003 COPYRIGHT (C) 2003 BIOLOGICAL ABSTRACTS INC.(R) FILE COVERS 1969 TO DATE. CAS REGISTRY NUMBERS AND CHEMICAL NAMES (CNs) PRESENT FROM JANUARY 1969 TO DATE. RECORDS LAST ADDED: 16 July 2003 (20030716/ED) => d all tot 179 L79 ANSWER 1 OF 2 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN **1998:386560** BIOSIS ΑN DN PREV199800386560 TΤ Dimerization of the polymeric immunoglobulin receptor controls its transcytotic trafficking. ΑU Singer, Karen L.; Mostov, Keith E. (1) (1) Dep. Anatomy, Univ. California, San Francisco, CA 94143-0452 USA CS SO Molecular Biology of the Cell, (April, 1998) Vol. 9, No. 4, pp. 901-915. ISSN: 1059-1524. DT Article LAEnglish Binding of dimeric immunoglobulin (Ig)A to the polymeric AB Ig receptor (pIgR) stimulates transcytosis of pigR across epithelial cells. Through the generation of a series of pIgR chimeric constructs, we have tested the ability of ligand to promote receptor dimerization and the subsequent role of receptor dimerization on its intracellular trafficking. Using the cytoplasmic domain of the T cell receptor-zeta chain as a sensitive indicator of receptor oligomerization, we show that a pIgR: zeta chimeric receptor expressed in Jurkat cells initiates a zeta-specific signal transduction cascade when exposed to dimeric or tetrameric IgA, but not when exposed to monomeric IgA. In

CC

BC

ΙT

ΙT

IT

L79

ΑN

DN

ΤI

ΑU

CS

SO

DT

LA

ΑB

BC

ΙT

IT

Miscellaneous Descriptors

addition, we replaced the pIgR's transmembrane domain with that of glycophorin A to force dimerization or with a mutant glycophorin transmembrane domain to prevent dimerization. Forcing dimerization stimulated transcytosis of the chimera, whereas preventing dimerization abolished ligand-stimulated transcytosis. We conclude that binding of dimeric IqA to the pIqR induces its dimerization and that this dimerization is necessary and sufficient to stimulate ${\tt pIgR}$ transcytosis. Biochemical Studies - General *10060 Cytology and Cytochemistry - Human *02508 Hominidae 86215 Major Concepts Biochemistry and Molecular Biophysics; Membranes (Cell Biology) Chemicals & Biochemicals dimeric immunoglobulin A; polymeric immunoglobulin receptor: dimerization, transcytotic trafficking; T cell receptor zeta chain cytoplasmic domain Miscellaneous Descriptors ligand-stimulated transcytosis ORGN Super Taxa Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia ORGN Organism Name Jurkat (Hominidae) ORGN Organism Superterms Animals; Chordates; Humans; Mammals; Primates; Vertebrates ANSWER 2 OF 2 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN **1994:156188** BIOSIS PREV199497169188 Protein traffic in polarized epithelial cells: The polymeric immunoglobulin receptor as a model system. Mostov, Keith Dep. Anat., Univ. Calif. San Francisco, San Francisco, CA 94143 USA Journal of Cell Science, (1993) Vol. 0, No. SUPPL. 17, pp. 21-26. ISSN: 0021-9533. General Review . English As a model system to study protein traffic in polarized epithelial cells, we have used the polymeric immunoglobulin receptor. This receptor travels first to the basolateral surface, where it can bind polymeric IgA or IgM. The receptor is then endocytosed and delivered to endosomes. The receptor is sorted into transcytotic vesicles, which are exocytosed at the apical surface. The 103-amino acid cytoplasmic domain of the receptor contains several sorting signals. The 17 residues closest to the membrane are an autonomous signal that is necessary and sufficient for basolateral sorting. For rapid endocytosis there are two independent signals, both of which contain critical tyrosine residues. Finally, transcytosis is signaled by phosphorylation of a particular serine. Cytology and Cytochemistry - Animal *02506 Biochemical Studies - Proteins, Peptides and Amino Acids 10064 Biochemical Studies - Carbohydrates 10068 Biophysics - Membrane Phenomena *10508 *12100 Movement Metabolism - Carbohydrates *13004 Metabolism - Proteins, Peptides and Amino Acids *13012 Immunology and Immunochemistry - General; Methods *34502 Animalia - Unspecified *33000 Major Concepts

Cell Biology; Membranes (Cell Biology); Metabolism; Physiology

IMMUNOGLOBULIN A; IMMUNOGLOBULIN M; PROTEIN SORTING; SORTING SIGNAL;

TRANSCYTOSIS

ORGN Super Taxa

Animalia - Unspecified: Animalia

ORGN Organism Name

animal (Animalia - Unspecified); Animalia (Animalia - Unspecified)

ORGN Organism Superterms

animals

=> d 182 bib ab tot

- L82 ANSWER 1 OF 25 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN
- AN 2002:164255 BIOSIS
- DN PREV200200164255
- TI Apical trafficking of wild-type and mutant forms of the polymeric immunoglobulin receptor.
- AU Low, Seng Hui (1); Mostov, Keith E.; Weimbs, Thomas
- CS (1) Department of Cell Biology, Lerner Research Institute, Cleveland Clinic Foundation, 9500 Euclid Avenue, NC10, Cleveland, OH, 44195 USA
- SO Molecular Biology of the Cell, (Dec., 2000) Vol. 11, No. Supplement, pp. 509a. http://www.molbiolcell.org/. print. Meeting Info.: 40th American Society for Cell Biology Annual Meeting San Francisco, CA, USA December 09-13, 2000 ISSN: 1059-1524.
- DT Conference
- LA English
- L82 ANSWER 2 OF 25 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN
- AN 2000:349921 BIOSIS
- DN PREV200000349921
- TI The polymeric immunoglobulin receptor mediates pneumococcal adherence and invasion across the human nasopharyngeal epithelial cells.
- AU Zhang, J. (1); Mostov, K.; Lamm, M.; Tuomanen, E. (1)
- CS (1) St. Jude Children's Research Hospital, Memphis, TN USA
- SO Abstracts of the General Meeting of the American Society for Microbiology, (2000) Vol. 100, pp. 71-72. print.

 Meeting Info.: 100th General Meeting of the American Society for Microbiology Los Angeles, California, USA May 21-25, 2000 American Society for Microbiology

 . ISSN: 1060-2011.
- DT Conference
- LA English
- SL English
- L82 ANSWER 3 OF 25 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN
- AN 1998:19938 BIOSIS
- DN PREV199800019938
- TI Role of tyrosine phosphorylation in **ligand**-induced regulation of transcytosis of the **polymeric immunoglobulin** receptor.
- AU Luton, Frederic; Cardone, Michael H.; Zhang, Min; Mostov, Keith E.
- CS Univ. Calif. San Francisco, Dep. Anat., San Francisco, CA 94143-0452 USA
- SO Molecular Biology of the Cell, (Nov., 1997) Vol. 8, No. SUPPL., pp. 88A.

 Meeting Info.: 37th Annual Meeting of the American Society for Cell

Meeting Info.: 37th Annual Meeting of the American Society for Cell Biology Washington, D.C., USA December 13-17, 1997 American Society for Cell Biology

- . ISSN: 1059-1524.
- DT Conference
- LA English
- L82 ANSWER 4 OF 25 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN

- 1997:384643 BIOSIS AN
- PREV199799683846 DN
- Evidence of dimerization of the polymeric immuno-globulin receptor upon TΙ binding to dIgA.
- Singer, K. L.; Mostov, K. E. ΑU
- Univ. California San Francisco, Dep. Anatomy, San Francisco, CA USA CS
- Journal of General Physiology, (1997) Vol. 110, No. 1, pp. 33A. SO Meeting Info.: Fifty-First Annual Meeting of the Society of General Physiologists Woods Hole, Massachusetts, USA September 4-6, 1997 ISSN: 0022-1295.
- Conference; Abstract; Conference DT
- LA English
- ANSWER 5 OF 25 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN L82
- AN 1996:442483 BIOSIS
- DN PREV199699164839
- Regulation of protein traffic in polarized epithelial cells: The TI polymeric immunoglobulin receptor model.
- Mostov, K. E. (1); Altschuler, Y.; Chapin, S. J. (1); ΑU Enrich, C.; Low, S.-H. (1); Luton, F.; Richman-Eisenstat, J.; Singer, K. L.; Tang, K.; Weimbs, T.
- (1) Dep. Anat., Univ. Calif., San Francisco, CA 94143-0452 USA CS
- COLD SPRING HARBOR LABORATORY.. Cold Spring Harbor Symposia on SO Quantitative Biology, (1995) Vol. 60, pp. 775-781. Cold Spring Harbor Symposia on Quantitative Biology; Protein kinesis: The dynamics of protein trafficking and stability. Publisher: Cold Spring Harbor Laboratory Press 10 Skyline Drive,

Plainview, New York 11803, USA.

Meeting Info.: Meeting Cold Spring Harbor, New York, USA 1995 ISSN: 0091-7451. ISBN: 0-87969-070-4 (paper), 0-87969-069-0 (cloth).

- DTBook; Conference
- English LA
- ANSWER 6 OF 25 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN L82
- 1996:54173 BIOSIS ΑN
- PREV199698626308 DN
- Dimerization of the polymeric immunoglobulin TΤ receptor using the transmembrane domain of the glycophorin: Effects of targeting.
- ΑU Singer, K. L.; Mostov, K. E.
- Dep. Anat., Univ. Calif., San Francisco, CA 94143 USA CS
- Molecular Biology of the Cell, (1995) Vol. 6, No. SUPPL., pp. 400A. SO Meeting Info.: Thirty-fifth Annual Meeting of the American Society for Cell Biology Washington, D.C., USA December 9-13, 1995 ISSN: 1059-1524.
- DΤ Conference
- LΑ English
- ANSWER 7 OF 25 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN L82
- ΑN 1995:52893 BIOSIS
- DN PREV199598067193
- TΙ Reconstitution of polymeric immunoglobulin receptor transcytosis in permeabilized MDCK cells.
- ΑU Apodaca, G.; Mostov, K. E.
- Dep. Anatomy, Univ. Calif., San Francisco, CA 94143 USA CS
- Molecular Biology of the Cell, (1994) Vol. 5, No. SUPPL., pp. 379A. SO Meeting Info.: Thirty-fourth Annual Meeting of the American Society for Cell Biology San Francisco, California, USA December 10-14, 1994 ISSN: 1059-1524.
- DT Conference
- LΑ English
- L82 ANSWER 8 OF 25 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN

- AN 1995:52635 BIOSIS
- DN PREV199598066935
- TI Interaction of calmodulin with the basolateral targeting signal of the polymeric immunoglobulin receptor.
- AU Chapin, S. J. (1); Enrich, C.; Aroeti, B. (1); Havel, R. J.; Mostov, K. E. (1)
- CS (1) Dep. Anat., Univ. Calif., San Francisco, CA 94143 USA
- SO Molecular Biology of the Cell, (1994) Vol. 5, No. SUPPL., pp. 334A. Meeting Info.: Thirty-fourth Annual Meeting of the American Society for Cell Biology San Francisco, California, USA December 10-14, 1994 ISSN: 1059-1524.
- DT Conference
- LA English
- L82 ANSWER 9 OF 25 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN
- AN 1995:52575 BIOSIS
- DN PREV199598066875
- TI Antisense to G-s inhibits transcytosis of dimeric IgA by the polymeric immunoglobulin receptor in Madin-Darby canine kidney cells.
- AU Okamoto, C. T.; Mostov, K. E.
- CS Dep. Anat. Cardiovascular Res. Inst., Univ. Calif., San Francisco, CA 94143-0452 USA
- SO Molecular Biology of the Cell, (1994) Vol. 5, No. SUPPL., pp. 323A. Meeting Info.: Thirty-fourth Annual Meeting of the American Society for Cell Biology San Francisco, California, USA December 10-14, 1994 ISSN: 1059-1524.
- DT Conference
- LA English
- L82 ANSWER 10 OF 25 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN
- AN 1995:51789 BIOSIS
- DN PREV199598066089
- TI IgA mediates IP3 production and protein kinase C activation in MDCK cells expressing the polymeric immunoglobulin receptor.
- AU Cardone, M. (1); Smith, B.; Mochly-Rosen, D.; Mostov, K. (1)
- CS (1) Dep. Anat. Biochem., Univ. California, San Francisco, CA 94143-0452 USA
- SO Molecular Biology of the Cell, (1994) Vol. 5, No. SUPPL., pp. 188A. Meeting Info.: Thirty-fourth Annual Meeting of the American Society for Cell Biology San Francisco, California, USA December 10-14, 1994 ISSN: 1059-1524.
- DT Conference
- LA English
- L82 ANSWER 11 OF 25 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN
- AN 1994:515737 BIOSIS
- DN PREV199497528737
- TI The polymeric immunoglobulin receptor is the major high affinity calmodulin binding protein in rat liver endosomes.
- AU Enrich, C. (1); Mostov, K. E.; Havel, J. R.
- CS (1) Dep. Biol. Cel. Anat. Patol., Fac. Med., Univ. Barcelona, Barcelona Spain
- SO Journal of Hepatology, (1994) Vol. 21, No. SUPPL. 1, pp. S75. Meeting Info.: 29th Annual Meeting of the European Association for the Study of the Liver Athens, Greece September 7-10, 1994 ISSN: 0168-8278.
- DT Conference
- LA English
- L82 ANSWER 12 OF 25 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN
- AN 1994:379395 BIOSIS

- DN PREV199497392395
- TI Both the G-s-alpha and beta-gamma subunits of the heterotrimeric G protein, G-S, control the sorting of the **polymeric** immunoglobulin receptor into transcytotic vesicles.
- AU Bomsel, Morgane (1); Mostov, Keith E.
- CS (1) Inst. Cochin de Genetique Moleculaire, 22 rue Mechain, 75014 Paris France
- SO Biochemical Society Transactions, (1994) Vol. 22, No. 2, pp. 463-468.

 Meeting Info.: 649th Meeting of the Biochemical Society London, England,
 UK December 19-21, 1993
 ISSN: 0300-5127.
- DT Conference
- LA English
- L82 ANSWER 13 OF 25 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN
- AN 1994:98961 BIOSIS
- DN PREV199497111961
- TI Internalization of the **polymeric immunoglobulin** receptor is decreased by mutation of a phosphorylated serine in its cytoplasmic domain.
- AU Okamoto, C. T. (1); Song, W.; Bomsel, M.; Mostov, K. E.
- CS (1) Dep. Anat., Univ. Calif., San Francisco, CA 94143-0452 USA
- SO Molecular Biology of the Cell, (1993) Vol. 4, No. SUPPL., pp. 437A. Meeting Info.: Thirty-third Annual Meeting of the American Society for Cell Biology New Orleans, Louisiana, USA December 11-15, 1993 ISSN: 1059-1524.
- DT Conference
- LA English
- L82 ANSWER 14 OF 25 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN
- AN 1994:98265 BIOSIS
- DN PREV199497111265
- TI Regulation of transcytosis of the **polymeric** immunoglobulin receptor by its physiological ligand.
- AU Song, W. (1); Bomsel, M.; Casanova, J.; Vaerman, J.-P.; Mostov, K.
- CS (1) Dep. Anat., Univ. Calif., San Francisco, CA 94143-0452 USA
- SO Molecular Biology of the Cell, (1993) Vol. 4, No. SUPPL., pp. 317A.

 Meeting Info.: Thirty-third Annual Meeting of the American Society for
 Cell Biology New Orleans, Louisiana, USA December 11-15, 1993
 ISSN: 1059-1524.
- DT Conference
- LA English
- L82 ANSWER 15 OF 25 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN
- AN 1994:97639 BIOSIS
- DN PREV199497110639
- TI Basolateral targeting of the **polymeric immunoglobulin** receptor from the trans-Golgi network and from basolateral endosomes of MDCK cells.
- AU Aroeti, B.; Kosen, P. A.; Kuntz, I. D.; Cohen, F. E.; Mostov, K. E.
- CS Dep. Anatomy, Univ. California, San Francisco, CA 94143 USA
- SO Molecular Biology of the Cell, (1993) Vol. 4, No. SUPPL., pp. 208A. Meeting Info.: Thirty-third Annual Meeting of the American Society for Cell Biology New Orleans, Louisiana, USA December 11-15, 1993 ISSN: 1059-1524.
- DT Conference
- LA English
- L82 ANSWER 16 OF 25 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN
- AN 1994:96951 BIOSIS
- DN PREV199497109951

- TI Phorbol ester mediated stimulation of transcytosis of the **polymeric immunoglobulin receptor** in MDCK cells involves protein kinase-C alpha translocation.
- AU Cardone, M. H. (1); Smith, Bradley L.; Mochly-Rosen, Daria; Mostov, K. E.
- CS (1) Dep. Anat., Univ. Calif., San Francisco, CA 94143-0452 USA
- SO Molecular Biology of the Cell, (1993) Vol. 4, No. SUPPL., pp. 90A. Meeting Info.: Thirty-third Annual Meeting of the American Society for Cell Biology New Orleans, Louisiana, USA December 11-15, 1993 ISSN: 1059-1524.
- DT Conference
- LA English
- L82 ANSWER 17 OF 25 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN
- AN 1993:263102 BIOSIS
- DN PREV199344125252
- TI Regulation of transcytosis of the **polymeric**immunoglobulin receptor in MDCK cells by protein kinase
 C.
- AU Cardonne, M. H.; Mostov, K. E.
- CS Univ. Calif., San Francisco, CA 94143-0452 USA
- SO Journal of Cellular Biochemistry Supplement, (1993) Vol. 0, No. 17 PART C, pp. 25.
 Meeting Info.: Keystone Symposium on Genetic and In Vitro Analysis of Cell Compartmentalization Taos, New Mexico, USA February 8-14, 1993
 ISSN: 0733-1959.
- DT Conference
- LA English
- L82 ANSWER 18 OF 25 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN
- AN 1993:241615 BIOSIS
- DN PREV199344114815
- TI Membrane traffic and transcytosis in polarized epithelial cells: Signals, mechanisms, and regulation.
- AU Mostov, K. (1); Apodaca, G.; Aroeti, B. (1); Song, W. (1); Bomsel, M.
- CS (1) Dep. Anat., Univ. Calif., San Francisco, CA 94143-0452 USA
- SO Journal of Cellular Biochemistry Supplement, (1993) Vol. 0, No. 17 PART C, pp. 49.

 Meeting Info.: Keystone Symposium on Emerging Principles for Vaccine

Meeting Info.: Keystone Symposium on Emerging Principles for Vaccine Development: Antigen Processing and Presentations Taos, New Mexico, USA February 8-14, 1993 ISSN: 0733-1959.

- DT Conference
- LA English
- L82 ANSWER 19 OF 25 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN
- AN 1992:65631 BIOSIS
- DN BR42:29531
- TI TRANSCYTOSIS OF PLACENTAL ALKALINE PHOSPHATASE POLYMERIC IMMUNOGLOBULIN RECEPTOR FUSIONS.
- AU APODACA G; MOSTOV K E
- CS DEP. ANATOMY, UNIVERSITY CALIFORNIA, SAN FRANCISCO, CALIF. 94143.
- ABSTRACTS OF PAPERS PRESENTED AT THE THIRTY-FIRST ANNUAL MEETING OF THE AMERICAN SOCIETY FOR CELL BIOLOGY, BOSTON, MASSACHUSETTS, USA, DECEMBER 8-12, 1991. J CELL BIOL. (1991) 115 (3 PART 2), 195A. CODEN: JCLBA3. ISSN: 0021-9525.
- DT Conference
- FS BR; OLD
- LA English
- L82 ANSWER 20 OF 25 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN
- AN 1990:190357 BIOSIS

- DN BR38:90680
- TI A SUBDOMAIN OF THE **POLYMERIC IMMUNOGLOBULIN**RECEPTOR CYTOPLASMIC TAIL SPECIFIES BASOLATERAL TARGETING IN MDCK
 CELLS.
- AU CASANOVA J E; MOSTOV K E
- CS DEP. ANAT., UNIV. CALIF. SAN FRANCISCO, SAN FRANCISCO, CALIF. 94143, USA.
- SO SYMPOSIUM ON GENETIC AND IN VITRO ANALYSIS OF CELL COMPARTMENTALIZATION HELD AT THE 19TH ANNUAL MEETINGS OF THE UNIVERSITY OF CALIFORNIA-LOS ANGELES SYMPOSIA ON MOLECULAR AND CELLULAR BIOLOGY, TAOS, NEW MEXICO, USA, FEBRUARY 3-9, 1990. J CELL BIOCHEM SUPPL. (1990) 0 (14 PART C), 38. CODEN: JCBSD7.
- DT Conference
- FS BR; OLD
- LA English
- L82 ANSWER 21 OF 25 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN
- AN 1990:59809 BIOSIS
- DN BR38:26229
- TI A SUBDOMAIN OF THE **POLYMERIC IMMUNOGLOBULIN RECEPTOR** CYTOPLASMIC TAIL SPECIFIES BASOLATERAL TARGETING IN MDCK CELLS.
- AU CASANOVA J E; MOSTOV K E
- CS DEP. ANATOMY, UC SAN FRANCISCO 94143.
- TWENTY-NINTH ANNUAL MEETING OF THE AMERICAN SOCIETY FOR CELL BIOLOGY, HOUSTON, TEXAS, USA, NOVEMBER 5-9, 1989. J CELL BIOL. (1989) 109 (4 PART 2), 295A.
 - CODEN: JCLBA3. ISSN: 0021-9525.
- DT Conference
- FS BR; OLD
- LA English
- L82 ANSWER 22 OF 25 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN
- AN 1989:171974 BIOSIS
- DN BR36:83215
- TI PHOSPHORYLATION AFFECTS POST-ENDOCYTIC SORTING OF THE POLYMERIC IMMUNOGLOBULIN RECEPTOR.
- AU CASANOVA J E; MOSTOV K E
- CS WHITEHEAD INST., CAMBRIDGE, MASS.
- SO JOINT MEETING OF THE AMERICAN SOCIETY FOR CELL BIOLOGY AND THE AMERICAN SOCIETY FOR BIOCHEMISTRY AND MOLECULAR BIOLOGY, SAN FRANCISCO, CALIFORNIA, USA, JANUARY 29-FEBRUARY 2, 1989. J CELL BIOL. (1988) 107 (6 PART 3), 447A.
 - CODEN: JCLBA3. ISSN: 0021-9525.
- DT Conference
- FS BR; OLD
- LA English
- L82 ANSWER 23 OF 25 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN
- AN 1989:171930 BIOSIS
- DN BR36:83171
- TI TRANSCYTOSIS AND SORTING OF THE POLYMERIC IMMUNOGLOBULIN RECEPTOR.
- AU MOSTOV K; CASANOVA J; BREITFELD P
- CS WHITEHEAD INST., CAMBRIDGE, MASS.
- SO JOINT MEETING OF THE AMERICAN SOCIETY FOR CELL BIOLOGY AND THE AMERICAN SOCIETY FOR BIOCHEMISTRY AND MOLECULAR BIOLOGY, SAN FRANCISCO, CALIFORNIA, USA, JANUARY 29-FEBRUARY 2, 1989. J CELL BIOL. (1988) 107 (6 PART 3), 439A.
 - CODEN: JCLBA3. ISSN: 0021-9525.
- DT Conference
- FS BR; OLD
- LA English

- L82 ANSWER 24 OF 25 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN
- AN 1987:157048 BIOSIS
- DN BR32:75175
- TI STRUCTURE AND FUNCTION OF THE RECEPTOR FOR POLYMERIC IMMUNOGLOBULINS.
- AU MOSTOV K E; FRIEDLANDER M; BLOBEL G
- CS WHITEHEAD INST., NINE CAMBRIDGE CENTER, CAMBRIDGE, MA 02142, USA.
- SO KAY, J., ET AL. (ED.). BIOCHEMICAL SOCIETY SYMPOSIA, NO. 51. GENES AND PROTEINS IN IMMUNITY; OXFORD, ENGLAND, JULY 1985. XIII+235P. THE BIOCHEMICAL SOCIETY: LONDON, ENGLAND. ILLUS. (1986) 0 (0), 113-116. CODEN: BSSYAT. ISSN: 0067-8694. ISBN: 0-904498-18-2.
- FS BR; OLD
- LA English
- L82 ANSWER 25 OF 25 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN
- AN 1987:99426 BIOSIS
- DN BR32:49227
- TI DELETION OF CYTOPLASMIC TAIL OF THE POLYMERIC IMMUNOGLOBULIN RECEPTOR PREVENTS BASOLATERAL LOCALIZATION AND ENDOCYTOSIS.
- AU MOSTOV K; DE BRUYN KOPS A; DEITCHER D
- CS WHITEHEAD INST., CAMBRIDGE, MA.
- SO TWENTY-SIXTH ANNUAL MEETING OF THE AMERICAN SOCIETY FOR CELL BIOLOGY, WASHINGTON, D.C., USA, DEC. 7-11, 1986. J CELL BIOL. (1986) 103 (5 PART 2), 8A.

 CODEN: JCLBA3. ISSN: 0021-9525.
- DT Conference
- FS BR; OLD
- LA English
- => fil wpix FILE 'WPIX' ENTERED AT 07:36:38 ON 23 JUL 2003 COPYRIGHT (C) 2003 THOMSON DERWENT
- FILE LAST UPDATED: 19 JUL 2003 <20030719/UP>
 MOST RECENT DERWENT UPDATE: 200346 <200346/DW>
 DERWENT WORLD PATENTS INDEX SUBSCRIBER FILE, COVERS 1963 TO DATE
- >>> NEW WEEKLY SDI FREQUENCY AVAILABLE --> see NEWS <<<
- >>> SLART (Simultaneous Left and Right Truncation) is now
 available in the /ABEX field. An additional search field
 /BIX is also provided which comprises both /BI and /ABEX <<</pre>
- >>> PATENT IMAGES AVAILABLE FOR PRINT AND DISPLAY <<<
- >>> FOR DETAILS OF THE PATENTS COVERED IN CURRENT UPDATES,
 SEE http://www.derwent.com/dwpi/updates/dwpicov/index.html <<<
- >>> FOR A COPY OF THE DERWENT WORLD PATENTS INDEX STN USER GUIDE,
 PLEASE VISIT:
 http://www.stn-international.de/training.center/patents/stn.guide:
- http://www.stn-international.de/training_center/patents/stn_guide.pdf <<<
- >>> FOR INFORMATION ON ALL DERWENT WORLD PATENTS INDEX USER
 GUIDES, PLEASE VISIT:
 http://www.derwent.com/userguides/dwpi_guide.html <<<</pre>

=> =>

=> d all abeq tech abex tot 1103

```
L103 ANSWER 1 OF 5 WPIX COPYRIGHT 2003 THOMSON DERWENT on STN
     2002-416628 [44]
                        WPIX
AN
DNC
    C2002-117522
     Complex useful for transporting active agent through epithelial barrier,
     has biologically active portion and target element directed to
     ligand that confers e.g. transcytotic properties to agent specific
     to ligand.
DC
     B04 D16
     BASU, A; CHAPIN, S; GLYNN, J M; HAWLEY, S; HOUSTON, L L;
IN
     SHERIDAN, P J
PΑ
     (ARIZ-N) ARIZEKE PHARM INC
CYC
    98
PΙ
    WO 2002028408 A2 20020411 (200244) * EN 379p
                                                     A61K038-00
       RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ
            NL OA PT SD SE SL SZ TR TZ UG ZW
        W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK
            DM DZ EC EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR
            KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PH PL PT RO
            RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW
    AU 2001096494 A 20020415 (200254)
                                                     A61K038-00
    EP 1324778
                   A2 20030709 (200345) EN
                                                     A61K047-48
         R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT
            RO SE SI TR
ADT WO 2002028408 A2 WO 2001-US30832 20011002; AU 2001096494 A AU 2001-96494
     20011002; EP 1324778 A2 EP 2001-977368 20011002, WO 2001-US30832 20011002
    AU 2001096494 A Based on WO 200228408; EP 1324778 A2 Based on WO 200228408
FDT
                     20010209; US 2000-237929P 20001002; US 2000-248478P
PRAI US 2001-267601P
     20001113; US 2000-248819P 20001114
     ICM A61K038-00; A61K047-48
IC
AΒ
     WO 200228408 A UPAB: 20020711
     NOVELTY - A complex or compound (I) comprising biologically active portion
     and a target element (II) directed to a ligand (L1) that confers
     transcellular, transcytotic or paracellular transporting properties to an
     agent specifically bound to L1, where (II) is not an antibody,
     is new. Alternatively, (I) comprises two or more (II) directed to one or
     more L1.
          DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the
     following:
          (1) a medical device or kit comprising (I);
          (2) a diagnostic composition (DC) comprising (I); and
          (3) a diagnostic kit comprising DC.
          ACTIVITY - None given.
         MECHANISM OF ACTION - None given.
          USE - (I) is useful for delivering a biologically active agent to an
     animal, for transporting an active agent through an epithelial or mucosal
     barrier, and for treating or identifying a disease in an animal (claimed).
     Dwg.0/26
FS
     CPI
FΑ
     AB; DCN
     CPI: B04-B01B; B04-C01; B04-D02; B04-E01; B04-G01; B04-H02; B04-H02A;
          B04-H02D; B04-H19; B04-J01; B04-J03A; B04-L01; B04-N02; B04-N03;
          B05-A03A; B05-A03B; B11-C07B; B12-K04E; D05-H09
TECH
                    UPTX: 20020711
     TECHNOLOGY FOCUS - BIOTECHNOLOGY - Preferred Complex: In (I), (II) is a
     nucleic acid or a polypeptide derived from a calmodulin, an AP-1 golgi
     adaptor or a bacterial polypeptide, and L1 is polyimmunoglobulin receptor
     (pIgR) stalk or a domain, conserved sequence or their region, or
     is a polypeptide having an amino acid sequence from LRKED, QLFVNEE, LNQLT,
     YWCKW, GWYWC, STLVPL, SYRTD and KRSSK, where L1 is in a region selected
     from R1 (from KRSSK to carboxy terminus of PIgR), R2a (from
     SYRTD to the carboxy terminus of {\bf PIgR}), R2b (from SYRTD to
     KRSSK), R3a (from STLVPL to the carboxy terminus of PIgR), R3b
     (from STLVPL to KRSSK), R3c (from STLVPL to SYRTD), R4a (from GWYWC to the
```

carboxy terminus of PIgR), R4b (from GWYWC to KRSSK), R4c (from GWYWC to SYRTD), R4d (GWYWC to STLVPL), R5a (from YWCKW to the carboxy terminus of PIgR), R5b (from YWCKW to KRSSK), R5c (from YWCKW to SYRTD), R5d (from YWCKW to STLVPL), R5e (from YWCKW to GWYWC), R6a (from LINQLT to the carboxy terminus to PIgR), R6b (from LNQLT to KRSSK), R6c (from LNQLT to SYRTD), R6d (from LNQLT to STLVPL), R6e (from LNQLT to GWYWC), R6f (from LNQLT to YWCKW), R7a (from QLFVNEE to the carboxy terminus of PIgR), R7b (from QLFVNEE to KRSSK), R7c (from QLFVNEE to SYRTD), R7d (from LNQLT to STLVPL), R7e (from QLFVNEE to GWYWC), R7f (from QLFVNEE to YWCKW), R7g (from QLFVNEE to LNQLT), R8a (from LRKED to the carboxy terminus of \mathbf{pIgR}), R8b (from LRKED to KRSSK), R8c (from LRKED to SYRTD), R8d (from LRKED to STLVPL), R8e (from LRKED to GWYWC), R8f (from LRKED to YWCKW), R8g (from LRKED to LNQLT) and R8h (from LRKED to QLFVNEE). (I) further comprises a biologically active portion that is not a targeting element. In (I), the compound further comprises a protein transduction domain (PTD) or membrane transport signals (MTS), where biologically active portion is a: (a) polypeptide including a peptidomimetic, nucleic acid, a lipid, a carbohydrate, a compound or complex comprising a metal which is from platinum(II), palladium(II), zinc and cobalt(III), a small molecule or their functional derivative, where the polypeptide is from growth factor, an interleukin, an immunogen, a hormone, an enzyme, an enzyme inhibitor, an antibody, a clotting factor, a receptor, a ligand for a receptor, a kinase, a phosphatase, a scaffold protein, an adaptor protein, a dominant negative mutant, a protease, a signaling molecule, a regulatory molecule, transporter, a transcriptional regulator, a nucleic acid binding protein or their functional derivatives, or is from insulin, interleukin (IL)-2, IL-4, human growth hormone (hGH), sCT and hCT; (b) a nucleic acid; or (c) second targeting element that is directed to a molecular target other than L1, which is preferably an antibody or its derivative, where the biologically active portion or its metabolite is absorbed from the lumen of an organ into the body of the animal, where lumen is from gastrointestinary, pulmonary, nasal, nasopharyngeal, pharyngeal, buccal, sublingual, vaginal, urogenital, ocular and tympanic lumen, ocular surface, uterine, urethral, bladder, mammary, salivary, lacrimal, respiratory sinus, biliary, sweat gland. (I) is delivered preferably to the blood, lymph, interstitial fluid or amniotic fluid of the animal or into the body with a pharmocokinetic profile that results in delivery of an effective dose of the compound or its active portion. (I) is capable of undergoing transcellular movement, baseolateral transcytosis, apical endocytosis, basolateral exocytosis, intracellular transport, and the complex or compound or its active portion is delivered to an intracellular compartment and is transported across the cellular barrier. In (I) comprising two or more (II), one of the (II) is identical or substantially identical, or different to one another (II). Preferably, (I) comprises n number of (II), where one or more of desirable attributes of the compound is enhanced as compared to a second compound having m targeting elements, where n and m are both whole integers, and n greater than m, where one or more desirable attributes is a change in affinity or avidity for L1, where a pharmacological property is from half-life, decreased secretion, efficacy and selectivity. (I) further comprises a detectable moiety. Preferred Composition: PC further comprises antiproteases or carrier polypeptides.

ABEX UPTX: 20020711

ADMINISTRATION - (I) is administered through oral, rectal (e.g. an enema or suppository) aerosol (e.g. for nasal or pulmonary delivery), parenteral or topical routes. Dosage of (I) 0.01-100, preferably 0.01-0.1 micro-g/kg. EXAMPLE - Single chain Fv antibody fragments (sFv) directed to epitopes in defined regions in polyimmunoglobulin receptor (pIgR) amino acid sequence was used in vitro genetic manipulation has been used to alter the reading frame of sFv5A to create derivatives that have

substitutions or insertions of amino acids with reactive sites. The template, pSyn expression vector encoding sFv5A, was amplified using primers 5'-AAATACCTATTGCCTACGGCAGCC-3' and 5'- ${\tt CGGAATTCCTACTAGCAGCCACCGCCACCTGCGGCCGCTAGGACGGTGACCTTGGTCCC-3'. \ \ \, The}$ polymerase chain reaction (PCR) product was cleaved with BamHI and EcoRI and ligated into expression vector DNA, where the resultant expression construct encoded sFv5A-G4Cys which had, from an amino-to carboxyterminal direction, a pelb leader sequence (for secretion in Escherichia coli encoded by vector sequences, sFv5A-Cys i.e. a heavy chain variable region, a spacer sequences (GGGGS repeated three times i.e. (G4S)3), a heavy chain variable region, another G4S linker, and a C-terminal cysteine residue that had been introduced into the sFv relative to sFv5A. Chemical conjugates of salmon calcitonin and sFv5AG4-Cys were prepared. Transcytosis assays were performed with sFv5A-G4Cys and with sFv5A-G4Cys-calcitonin conjugates. The transcytosis assays with sFv5A-G4Cys showed that the sFv5A-G4Cys preparation was a mixture of monomers and dimers. A portion of the sFv preparation migrates as a dimer on non-reducing sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE). The dimer species was probably produced by covalent or non-covalent interactions that occurred prior to boiling in SDS. Thus, by comparing the monomer and dimer bands on the gel, one can monitor monomer and dimer transcytosis in the same sample. Transcytosis of sFv5A-G4Cys dimers was typically greater than 10 %, whereas transcytosis sFv5A-G4Cys monomers was usually less than 10 often less than 5 %. The transcytosis assays with sFv5A-G4Cys showed that the preparation of monomer Fv5A-G4Cys-calcitonin that was tested shows 2 conjugate species on SDS-PAGE. These species of conjugates behave differently. Transcytosis of the gel-monomer conjugate was relatively inefficient, resembling that of the sFv5A-G4Cys monomer. In contrast, transcytosis of the gel-dimer conjugate was relatively efficient.

```
L103 ANSWER 2 OF 5 WPIX COPYRIGHT 2003 THOMSON DERWENT on STN
AN
     2001-611619 [70]
                       WPIX
    C2001-182806
DNC
     New ligands binding to a specific region of a polymeric
TΙ
     immunoglobulin receptor, useful for transporting
     therapeutic or diagnostic compositions into or across cells expressing
    pIgR e.g. in drug delivery.
DC
     B04 D16
     CHAPIN, S J; MOSTOV, K E; RICHMAN-EISENSTAT, J
IN
     (REGC) UNIV CALIFORNIA; (CHAP-I) CHAPIN S J; (MOST-I) MOSTOV K E; (RICH-I)
PA
     RICHMAN-EISENSTAT J
CYC
     WO 2001072846 A2 20011004 (200170) * EN 102p
                                                     C07K016-28
PΙ
        RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ
            NL OA PT SD SE SL SZ TR TZ UG ZW
         W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK
            DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ
            LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD
            SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW
     AU 2001052970 A 20011008 (200208)
                                                     C07K016-28
                                                                      <--
     US 2002102657 A1 20020801 (200253)
                                                     C12P021-04
                   A2 20030102 (200310) EN
                                                     C07K016-28
     EP 1268555
         R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT
            RO SE SI TR
ADT WO 2001072846 A2 WO 2001-US9699 20010326; AU 2001052970 A AU 2001-52970
     20010326; US 2002102657 A1 Provisional US 2000-192197P 20000327,
     Provisional US 2000-192198P 20000327, US 2001-818247 20010326; EP 1268555
```

FDT AU 2001052970 A Based on WO 200172846; EP 1268555 A2 Based on WO 200172846

PRAI US 2000-192198P 20000327; US 2000-192197P 20000327; US 2001-818247

A2 EP 2001-926437 20010326, WO 2001-US9699 20010326

20010326

TC

ICM C07K016-28; C12P021-04

ICS A61K031-00; A61K031-7088; A61K038-00; **A61K039-395**;
A61K047-48; A61K048-00; A61P011-00; C07K019-00; C12N005-06

ICA C07K014-705

AB

WO 200172846 A UPAB: 20011129

NOVELTY - Ligands that bind specifically to a region of an animal cell polymeric immunoglobulin receptor (pIgR) are new. The pIgR cleaves to produce a stalk region remaining attached to the cell and a secretory component existing in the organ of interest in several forms. The ligands do not bind to the stalk or the most abundant form of SC present in the organ under physiological conditions.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) a **ligand** as above in which the **ligand** does not substantially bind to a peptide comprising 31 amino acids that are cell-membrane-proximal to the initial cleavage site;
- (2) a **ligand** as above in which **ligand** binding reduces proteolytic cleavage of SC by at least one-third compared to SC cleavage from cells without **ligand** binding;
- (3) introducing the novel ligand or the ligand of (1) or (2) into a cell (optionally an epithelial cell) of an animal organ, by binding ligand to pIgR;
- (4) a conjugate fusion protein or complex comprising a ligand as in (2) and a biologically active component;
- (5) attaching a **ligand** of (2), or conjugate fusion protein or complex of (4), to a cell expressing **pIgR**, by binding **ligand** to receptor, optionally in which **ligand** is internalized into the cell after binding; and
- (6) transcytosing a **ligand** from an apical to a basolateral side of a cell expressing **pIgR** of an animal organ, by binding the novel **ligand** or the **ligand** of (1) or (2) to **pIgR**.

USE - The ligands are useful for transporting therapeutic or diagnostic compositions into or across cells expressing pIgR, useful to introduce or transport ligands such as antibodies and/or to deliver biologically active components such as proteins, nucleic acids or detectable labels. They are used to deliver therapeutic compositions to mucosal surfaces such as the gastro-intestinal tract, respiratory system etc. in humans. They are also useful to label cells expressing pIgR, e.g. to distinguish epithelial cells from a mixed cell population in pathology studies or to aid in carcinoma diagnosis (since pIgR expression is reduced in carcinomas relative to normal epithelium). They can also be used to deliver veterinary compositions, especially in mammals such as farm, domestic or wild mammals or birds e.g. birds reared for human consumption.

Dwg.0/5

FS CPI

FA AB; DCN

MC CPI: B04-C01B; B04-C01C; B04-G01; B04-N04; B11-C07A; B12-K04A; D05-H09; D05-H11

TECH UPTX: 20011129

TECHNOLOGY FOCUS - BIOTECHNOLOGY - Preferred Ligands: The ligand is an antibody, particularly a humanized antibody, and the animal is a bird or a mammal, especially a human. The organ of interest is preferably a lung, small intestine, large intestine, liver-biliary tree, stomach, salivary gland, vagina, lacrimal gland, uterus, mammary gland, nasal passage, or sinus. The ligand may optionally further comprise a biologically active component (e.g. a polynucleotide, protein, radioisotope, lipid, carbohydrate, peptidomimetric, antiinfective, antibiotic, or small molecule), especially when the organ is a lung and the component is a polynucleotide encoding the wildtype cystic fibrosis transmembrane conductance regulator. Preferred Methods: Introducing a ligand into a cell of (3) uses

a ligand of (2). The rate of internalization of a first ligand binding to SC can be increased in cells secreting pIgR from an apical surface by binding pIgR to ligand of (2), and binding first ligand to the SC. The rate of transcytosis a first ligand binding to SC from an apical to a basolateral side of a cell can be increased in animal cells secreting pIgR by binding pIgR at the cell apical side to ligand of (2), and binding first ligand to the SC.

ABEX

UPTX: 20011129

SPECIFIC SEQUENCES - Ligands binding to epitope sequences (I)-(VII) are specifically claimed. Also claimed are ligands binding to one of 30 26-131 residue peptides derived from human pIgR (all fully defined in the specification). GlnAspProArgLeuPhe (I) LeuAspProPheLeuPhe (II) LysAlaIleGlnAspProArgLeuPhe (III); LeuAspProArgLeuPheAlaAspGluArgIle (IV); AspGluAsnLysAlaAsnLeuAspProArgLeuPhe (V) ArgLeuPheAlaAspGluArgGluIle (VI); LeuAspProArgLeuPheAlaAspGlu (VII).

EXAMPLE - A Fab fragment reactive to the B region of human pIgR was produced routinely and linked to poly (L-lysine) as previously described (Ferkol et al., J. Clin. Invest., 92:2394-2400 (1993)). A plasmid containing the wildtype cystic fibrosis transconductance regulator (CFTR) gene was ligated to a cytomegalovirus early promoter and inserted into pCB6. Plasmid DNA was combined with Fab-polylysine in 3M NaCl to produce Fab-polylysine-DNA complex. Complex was dissolved in 0.1 ml phosphate buffered saline and 100 micro-l applied into nares of anesthetized, pathogen-free Sprague-Dawley rats to target the CFTR gene into cells expressing pIgR. CFTR transcription was assayed by immunofluorescence assay of CFTR protein; no results are given.

L103 ANSWER 3 OF 5 WPIX COPYRIGHT 2003 THOMSON DERWENT on STN 2000-665249 [64] WPIX AN DNN N2000-493019 DNC C2000-201579 Quantitatively detecting ligand movement across a biological TТ membrane, comprises contacting assay-compatible infrared fluorescent labeled ligands with a receptor. B04 D16 E23 E24 S03 DC ALTSCHULER, Y; MOSTOV, K IN PA (REGC) UNIV CALIFORNIA CYC C12Q001-00 PΙ WO 2000063418 A1 20001026 (200064)* EN 23p RW: AT BE CH CY DE DK ES FI FR GB GR IE IT LU MC NL PT SE W: AU CA JP AU 2000042455 A 20001102 (200107) C120001-00 WO 2000063418 A1 WO 2000-US10173 20000414; AU 2000042455 A AU 2000-42455 ADT 20000414 FDT AU 2000042455 A Based on WO 200063418 PRAI US 1999-292274 19990415 ICM C120001-00 TC C07H019-20; C12Q001-02; C12Q001-04; C12Q001-32; G01N033-00; ICS G01N033-53 WO 200063418 A UPAB: 20001209 ΑB NOVELTY - Quantitatively detecting ligand movement across a biological membrane, comprising contacting a ligand comprising an assay-compatible infrared fluorescent label with a receptor, where the receptor binds and transports the ligand across a biological membrane, and quantitatively detecting fluorescence to indicate. ligand transport, is new.

(a) contacting a ligand comprising a compatible fluorescent

identifying an agent which modulates movement of a ligand across

a membrane, comprising:

DETAILED DESCRIPTION - An INDEPENDENT CLAIM is also included for

label with a receptor, in the presence of a candidate agent, under conditions where in the absence of the agent the receptor transports a first amount of the **ligand** across a membrane;

- (b) quantitatively detecting fluorescence as an indicator of a second amount of the **ligand** transported across the membrane; and
- (c) comparing the two amounts of transported **ligand**, a difference indicates that the agent modulates movement of the **ligand** across the membrane.

USE - For detecting the movement of macromolecules across biological membranes, and for identifying modulators of the molecular transport (claimed). The macromolecules may be e.g. hormones, cytokines, antibiotics, cytotoxins, chemokines, chemotactic factors, growth factors or neurotransmitters.

ADVANTAGE - The novel method uses infrared labels which are sensitive enough to replace the most sensitive existing labels, such as radiolabels and fluorescent labels, and do not interfere with the transport process. Radiolabels are unsuitable for high-throughput application, and fluorescent label use is limited by spectroscopic interference, the infrared labels overcome these problems.

Dwg.0/0

FS CPI EPI

FA AB; GI; DCN

MC CPI: B04-H01; B04-J01; B06-D01; B06-D13; B06-D18; B08-D01; B11-C07B3;

B12-K04; D05-H09; E23; E24-A03

EPI: S03-E14H4

TECH

UPTX: 20001209

TECHNOLOGY FOCUS - BIOTECHNOLOGY - Preferred Method: The membrane comprises a plasma or endosomal membrane and the transport across the membrane is by endocytosis, exocytosis or translocation. The membrane may alternatively comprise a layer of cells and the transport across the membrane is by transcytosis. The ligand is a protein, e.g. immunoglobulin (Ig)A or transferrin, and the detecting step uses a scanning fluorimeter. The method is repeated in massive parallel in distinct elements of an assay array, preferably distinct wells of a multiwell plate. The label comprises a dye having formula (I). R1, R6, R7 and R12 = independently, substituted or unsubstituted V elements, substituted or unsubstituted VI elements, or substituted or unsubstituted alkyl, alkenyl, alkynyl, aryl, heteroaryl, heteroalkyl, heteroaryl, and acyl substituents; R2-R5 and R8-R11 = independently, substituted or unsubstituted alkyl, alkenyl, alkynyl, aryl, heteroalkyl, heteroaryl, and acyl substituents; and

R13 = substituted or unsubstituted aryloxy or heteroaryloxy.

ABEX UPTX: 20001209
EXAMPLE - Madin-Darby canine

EXAMPLE - Madin-Darby canine kidney epithelial cells were cultured transfected in permeable filter supports to form a well polarized monolayer, essentially reconstituting a simple epithelial tissue, with an apical surface in contact with the overlying medium. Material added to the medium underneath the filter can diffuse through the filter to reach the basolateral surface. For immunoglobulin (Ig)A transport, the cells were transfected with cDNA for rabbit pIgR, and the exogenously expressed pigR functions as in vivo. IgA is labeled and added to the basolateral medium. Endocytosis was allowed to proceed for 10 minutes at 37 degrees C, and the cells were washed extensively during a 5 minute period. Finally, the release of IgA into the apical medium, or onto the basolateral medium was followed by sampling the medium over a 2 hour period. Detection of transcytosed IgA labeled with several infrared dyes was easily accomplished, by using 0.3 micro-g infrared fluorescent IgA. Only 0.27 % of the total apical medium was spotted on the filter, but the very small amount of transcytosed IgA was detected on the apical side of the cells. The system was found to be 170-280 fold more sensitive than radio-iodination

```
L103 ANSWER 4 OF 5 WPIX COPYRIGHT 2003 THOMSON DERWENT on STN
ΑN
     2000-549134 [50]
                       WPIX
    C2000-163952
DNC
     Novel polypeptides containing plgR-binding domains used for
ΤI
     targeting and transport to the mucosal epithelia, in the treatment of
     disorders accessible to the mucosal epithelia, e.g. asthma.
DC
     B04 D16
     CAPRA, J D; HEXHAM, J M; MANDECKI, W; WHITE, K
ΙN
     (DGIB-N) DGI BIOTECHNOLOGIES; (OKLA-N) OKLAHOMA MEDICAL RES FOUND; (TEXA)
PΑ
     UNIV TEXAS SYSTEM; (DGIB-N) DGI BIOTECHNOLOGIES INC
CYC
     WO 2000047611 A2 20000817 (200050)* EN 137p
                                                     C07K014-00
PI
        RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL
            OA PT SD SE SL SZ TZ UG ZW
        W: AE AL AM AT AU AZ BA BB BG BR BY CA CH CN CR CU CZ DE DK DM EE ES
            FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS
            LT LU LV MA MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL
            TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW
     AU 2000027597 A 20000829 (200062)
                                                     C07K014-00
     EP 1151000
                   A2 20011107 (200168) EN
                                                     C07K014-00
         R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT
            RO SE SI
     JP 2002539771 W 20021126 (200307)
                                             166p
                                                     C12N015-09
    WO 2000047611 A2 WO 2000-US3650 20000211; AU 2000027597 A AU 2000-27597
ADT
     20000211; EP 1151000 A2 EP 2000-906030 20000211, WO 2000-US3650 20000211;
     JP 2002539771 W JP 2000-598527 20000211, WO 2000-US3650 20000211
    AU 2000027597 A Based on WO 200047611; EP 1151000 A2 Based on WO
     200047611; JP 2002539771 W Based on WO 200047611
PRAI US 1999-119932P 19990212
     ICM C07K014-00; C12N015-09
         A61K038-00; A61K045-00; A61K047-48; A61P001-00; A61P001-12;
         A61P001-14; A61P001-16; A61P001-18; A61P011-00; A61P011-06;
          A61P013-12; A61P029-00; A61P031-00; A61P031-04; A61P031-10;
          A61P031-14; A61P031-16; A61P031-18; A61P031-20; A61P031-22;
          A61P033-02; A61P035-00; C07K014-47; C07K014-705;
          C07K019-00; C12N015-10; C12N015-62
     WO 200047611 A UPAB: 20001010
AΒ
     NOVELTY - A 10-50 residue peptide (I) comprising a pIgR-binding
     domain, is new.
          DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the
          (1) a fusion protein comprising a pIgR-binding domain
     covalently linked to a non-antibody peptide or polypeptide;
          (2) a polynucleotide encoding the fusion protein of (1);
          (3) targeting an agent to a mucosal epithelium comprising
     administering to a mammal a targeting complex comprising the agent and
     (I), the complex binds to, and is taken up by, cells expressing
     pIgR, and is transported to the mucosal epithelium;
          (4) targeting a non-antibody peptide or polypeptide to a
     mucosal epithelium, comprising administering the fusion protein of (1) to
     a mammal, the protein binds to, and is taken up by, cell expressing
     pIgR, and is transported to the mucosal epithelium;
          (5) delivering an agent to a cell, comprising contacting (I) with a
     cell expressing pIgR; and
          (6) delivering a non-antibody peptide or polypeptide to a
   . cell, comprising contacting the fusion protein of (1) with a cell
     expressing plqR.
          ACTIVITY - Antiasthmatic; antiinflammatory; antiinfectious;
     cytostatic; antiulcer; antidiarrheal; hepatropic; virucide; vasotropic;
     anti-human immunodeficiency virus; antibacterial. No biological data is
```

USE - For targeting and transport to the mucosal epithelium

given.

MECHANISM OF ACTION - None given.

(claimed), for the prevention or treatment of diseases, ailments or conditions that are accessible to mucosal epithelia, including asthma, bronchitis, emphysema, cystic fibrosis, bronchiectasis, bronchiolitis, pulmonary edema, viral tracheobronchitis, sleep apnea syndrome, infectious diseases, neoplastic conditions, Loffler's syndrome, kyphocliosis, dysphagia, peptic ulcers, diarrheal diseases, ulcerative colitis, Crohn's disease, hepatitis, cirrhosis, hemorrhoids, systemic vasculitis, acquired immunodeficiency syndrome, gonorrhea, syphilis and chlamydia. (I) can be attached to a detectable label for use in diagnostics.

FS CPI

FA AB; DCN

MC CPI: B04-C01; B04-E02; B04-E03; B04-N04; B04-N04A; B12-K04A; B14-A01; B14-E02; B14-E04; B14-E08; B14-E10C; B14-F02; B14-G01B; B14-H01; B14-K01; B14-N07C; B14-N12; D05-C11; D05-H09; D05-H12C; D05-H17C

UPTX: 20001010

TECH

TECHNOLOGY FOCUS - ORGANIC CHEMISTRY - Preferred Peptide: (I) is 10, 15, 20, 25, 30, 35, 40, 45 or 50 residues in length, and further comprises a linking moiety, preferably SMTP, SPDP, LC-SPDP, Sulfo-LC-SDPD, SMCC, Sulfo-SMCC, MBS, Sulfo-MBS, SIAB, Sulfo-SIAB, SMPM, Sulfo-SMPB, EDC/Sulfo-NHS, or ABH, attached to the peptide. The linking moiety may be further attached to an agent, preferably a peptide, polypeptide, oligonucleotide, polynucleotide, detachable label or drug. The polypeptide is an enzyme, antibody region, region mediating protein-protein interaction, cytokine, growth factor, hormone, toxin, tumor suppressor, transcription factor, or apoptosis inducer. The polynucleotide encodes a polypeptide, a single chain antibody, an antisense construct or a ribozyme. The detectable label is rhodamine, fluorescein, green fluorescent protein or a radiolabel. The drug is an antibiotic, DNA damaging agent, enzyme inhibitor, or metabolite. Alternatively, (I) further comprises a non-pigR targeting agent linked to the peptide. The targeting agent is an antigen binding domain of an antibody, a receptor ligand or ligand binding domain. (I) may comprise two pIgR-binding domains, and further comprise the linking agent and the agent.

Comprise the linking agent and the agent.

Preferred Fusion Protein: The domain is Calpha3 domain. The nonantibody peptide or polypeptide is an enzyme, antibody
region, region mediating protein-protein interaction, cytokine, growth
factor, hormone, toxin, tumor suppressor, transcription factor, or
apoptosis inducer.

Preferred Complex: The targeting complex further comprises a non-pigR targeting agent.

TECHNOLOGY FOCUS - BIOTECHNOLOGY - Preferred Method: Prior to performing the method of (5), the cell is transformed with an expression construct encoding pigR under the control of a promoter.

Preparation: (I) can be produced by standard recombinant techniques. ABEX UPTX: 20001010

SPECIFIC POLYPEPTIDES - (I) comprises one of 41 polypeptide sequences containing 9-45 residues, all fully defined in the specification, e.g. GlnGluProSerGlnGlyThrThrThr, ArgGlyGlyAsnGlyAlaLeuSerTrpArgGlyPheGlyTrpAla HisAspSerTrpPheProTrpPheGlyGly, and GlyTrpLeuGlyGluGlyTrpTrpGluLeuLeu (claimed).

ADMINISTRATION - The mucosal epithelium targeting complex is administered by oral, inhalation, ocular, nasal, vaginal, rectal, intravenous, subcutaneous, intramuscular, or intraarterial routes.

EXAMPLE - No relevant examples are given.

L103 ANSWER 5 OF 5 WPIX COPYRIGHT 2003 THOMSON DERWENT on STN AN 1998-042123 [04] WPIX DNC C1998-014108

```
ΤI
     Ligand that binds the stalk of a cell's polymeric
     immunoglobulin receptor - useful to target to, into or
     across mammalian epithelial cell biologically active component, e.g.
     nucleic acid, protein, lipid, carbohydrate, etc.
DC
    MOSTOV, K E; RICHMAN-EISENSTAT, J; MOSTOV, K
ΙN
     (REGC) UNIV CALIFORNIA
PΑ
CYC
    77
                   A1 19971211 (199804) * EN
                                                     C07K016-00
     WO 9746588
                                              42p
PΙ
        RW: AT BE CH DE DK EA ES FI FR GB GH GR IE IT KE LS LU MC MW NL OA PT
            SD SE SZ UG
        · W: AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI GB GE
            HU IL IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG MK MN MW MX
            NO NZ PL PT RO RU SD SE SG SI SK TJ TM TR TT UA UG UZ VN YU
                   A 19980105 (199821)
                                                     C07K016-00
     AU 9730632
     EP 934338
                   A1 19990811 (199936)
                                         ΕN
                                                     C07K016-00
         R: AT BE CH DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE
                  A 19990630 (199944)
     CN 1221428
                                                     C07K016-00
     US 6042833
                   Α
                     20000328 (200023)
                                                     A61K038-16
     JP 2000511432 W 20000905 (200047)
                                              46p
                                                     C12N015-09
                   B 20010111 (200108)
                                                     C07K016-00
     AU 728587
     IL 127238
                   A 20010724 (200147)
                                                     C07K016-00
     US 6340743
                   B1 20020122 (200208)
                                                     C07K016-28
     RU 2191781
                   C2 20021027 (200281)
                                                     C07K016-42
    WO 9746588 A1 WO 1997-US7944 19970514; AU 9730632 A AU 1997-30632
ADT
     19970514; EP 934338 A1 EP 1997-925515 19970514, WO 1997-US7944 19970514;
     CN 1221428 A CN 1997-195238 19970514; US 6042833 A Provisional US
     1996-18958P 19960604, US 1997-856383 19970514; JP 2000511432 W WO
     1997-US7944 19970514, JP 1998-500584 19970514; AU 728587 B AU 1997-30632
     19970514; IL 127238 A IL 1997-127238 19970514; US 6340743 B1 Provisional
     US 1996-18958P 19960604, Div ex US 1997-856383 19970514, US 1999-475088
     19991230; RU 2191781 C2 WO 1997-US7944 19970514, RU 1999-100279 19970514
FDT AU 9730632 A Based on WO 9746588; EP 934338 A1 Based on WO 9746588; JP
     2000511432 W Based on WO 9746588; AU 728587 B Previous Publ. AU 9730632,
     Based on WO 9746588; US 6340743 B1 Div ex US 6042833; RU 2191781 C2 Based
     on WO 9746588
                      19960604; US 1997-856383
PRAI US 1996-18958P
                                                 19970514; US 1999-475088
     19991230
     ICM A61K038-16; C07K016-00; C07K016-28; C07K016-42; C12N015-09
TC
         A61K039-385; A61K039-395; C07K016-46; C12N015-13
     ICS
AB
          9746588 A UPAB: 19980126
     WO
       Ligand that specifically binds the stalk of a polymeric
     immunoglobulin receptor (pIgR) of a cell, but
     not the secretory component of plgR under
     physiological conditions, is claimed.
          USE - The ligand, which can be introduced into a cell
     expressing a pIgR by attaching to the stalk of the pIgR
     , can be used to target to, into or across the apical or basolateral
     surface of a mammalian epithelial cell, a biologically active component
     selected from a nucleic acid (preferably encoding the wild type cystic
     fibrosis transmembrane conductance regulator), protein, radioisotope,
     lipid or carbohydrate, e.g. an anti-inflammatory, antisense
     oligonucleotide, antibiotic or anti-infective.
     Dwg.0/0
     CPI
FS
FΑ
     AΒ
     CPI: B04-G21; B04-G22; B04-N02A; B11-C07A; B12-K04
MC
=> fil dpci
```

FILE 'DPCI' ENTERED AT 07:41:21 ON 23 JUL 2003

COPYRIGHT (C) 2003 THOMSON DERWENT

FILE LAST UPDATED: 21 JUL 2003 <20030721/UP>
PATENTS CITATION INDEX, COVERS 1973 TO DATE

```
>>> LEARNING FILE LDPCI AVAILABLE <<<
```

=> d all tot 1113

L113 ANSWER 1 OF 3 DPCI COPYRIGHT 2003 THOMSON DERWENT on STN

AN 2001-611619 [70] DPCI

DNC C2001-182806

New ligands binding to a specific region of a polymeric immunoglobulin receptor, useful for transporting therapeutic or diagnostic compositions into or across cells expressing pIgR e.g. in drug delivery.

DC B04 D16

IN CHAPIN, S J; MOSTOV, K E; RICHMAN-EISENSTAT, J

PA (REGC) UNIV CALIFORNIA; (CHAP-I) CHAPIN S J; (MOST-I) MOSTOV K E; (RICH-I) RICHMAN-EISENSTAT J

CYC 96

PI WO 2001072846 A2 20011004 (200170)* EN 102p C07K016-28 <-RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ
NL OA PT SD SE SL SZ TR TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

AU 2001052970 A 20011008 (200208) C07K016-28 <-US 2002102657 A1 20020801 (200253) C12P021-04 <-EP 1268555 A2 20030102 (200310) EN C07K016-28 <--

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI TR

ADT WO 2001072846 A2 WO 2001-US9699 20010326; AU 2001052970 A AU 2001-52970 20010326; US 2002102657 A1 Provisional US 2000-192197P 20000327, Provisional US 2000-192198P 20000327, US 2001-818247 20010326; EP 1268555 A2 EP 2001-926437 20010326, WO 2001-US9699 20010326

FDT AU 2001052970 A Based on WO 200172846; EP 1268555 A2 Based on WO 200172846 PRAI US 2000-192198P 20000327; US 2000-192197P 20000327; US 2001-818247 20010326

IC ICM C07K016-28; C12P021-04 ICS A61K031-00; A61K031-7088; A61K038-00; A61K039-395; A61K047-48; A61K048-00; A61P011-00; C07K019-00; C12N005-06

ICA · C07K014-705

FS CPI

CTCS CITATION COUNTERS

PNC.DI 0	Cited Patents Count (by inventor)
PNC.DX 3	Cited Patents Count (by examiner)
IAC.DI 0	Cited Issuing Authority Count (by inventor)
IAC.DX 2	Cited Issuing Authority Count (by examiner)
PNC.GI 0	Citing Patents Count (by inventor)
PNC.GX 0	Citing Patents Count (by examiner)
IAC.GI 0	Citing Issuing Authority Count (by inventor)
IAC.GX 0	Citing Issuing Authority Count (by examiner)
CRC.I 0 CRC.X 4	Cited Literature References Count (by inventor) Cited Literature References Count (by examiner)

CDP CITED PATENTS

UPD: 20030627

Cited by Examiner

CITING PATENT CAT CITED PATENT ACCNO _____ US 5972900 A 1995-351156/45 WO 200172846 A A PA: (UYCA-N) UNIV CASE WESTERN RESERVE; (UYOH-N) UNIV OHIO; (OHIS) UNIV OHIO STATE; (OHIS) UNIV OHIO IN: FERKOL, T W; HANSON, R W; PERALES, J C; DAVIS, P B; ZIADY, A A 1996-333987/33 WO 9621012 (PLAN-N) PLANET BIOTECHNOLOGY INC; (UNME-N) UNITED PA: MEDICAL & DENTAL SCHOOLS GUYS; (PLAN-N) PLANT BIOTECHNOLOGY INC; (SCRI) SCRIPPS RES INST; (HIAT-I) HIATT A C; (LEHN-I) LEHNER T; (MAJK-I) MA J K -; (MOST-I) MOSTOV K E IN: HIATT, A C; MA, J K; LEHNER, T; MA, J K C; HEIN, M B; MOSTOV, K E; MA, J K -WO 9746588 A 1998-042123/04 PA: (REGC) UNIV CALIFORNIA IN: MOSTOV, K E; RICHMAN-EISENSTAT, J; MOSTOV, K

REN LITERATURE CITATIONS UPR: 20030627

Citations by Examiner

CITING PATENT	CAT	CITED LITERATURE
WO 200172846	A	E. ECKMAN ET AL.: "In vitro transport of active alphal-antitrypsin to the apical surface of epithelia by targeting the polymeric immunoglobulin receptor." AMERICAN JOURNAL OF RESPIRATORY CELL AND MOLECULAR BIOLOGY, vol. 21, no. 2, August 1999 (1999-08), pages 246-252, XP001031177 New York, NY, USA
WO 200172846	A	P. KRAJCI ET AL.: "Molecular cloning and exon-intron mapping of the gene encoding human transmembrane secretory component (the poly-Ig receptor)." EUROPEAN JOURNAL OF IMMUNOLOGY, vol. 22, no. 9, September 1992 (1992-09), pages 2309-2315, XP000567240 Weinheim, Germany
WO 200172846	A .	K. MOSTOV: "Transepithelial transport of immunoglobulins." ANNUAL REVIEW OF IMMUNOLOGY, vol. 12, 1994, pages 63-84, XP001053221 Palo Alto, CA, USA
WO 200172846	A .	T. FERKOL ET AL.: "Gene transfer into respiratory epithelial cells by targeting the polymeric immunoglobulin receptor." JOURNAL OF CLINICAL INVESTIGATION, vol. 92, no. 5, November 1993 (1993-11), pages 2394-2400, XP001053217 New York, NY, USA

L113 ANSWER 2 OF 3 DPCI COPYRIGHT 2003 THOMSON DERWENT on STN

AN 2000-665249 [64] DPCI

DNN N2000-493019 DNC C2000-201579

TI Quantitatively detecting ligand movement across a biological membrane, comprises contacting assay-compatible infrared fluorescent labeled ligands with a receptor.

DC B04 D16 E23 E24 S03

IN ALTSCHULER, Y; MOSTOV, K

PA (REGC) UNIV CALIFORNIA

CYC 21

PI WO 2000063418 A1 20001026 (200064) * EN 23p C12Q001-00

```
RW: AT BE CH CY DE DK ES FI FR GB GR IE IT LU MC NL PT SE
        W: AU CA JP
    AU 2000042455 A 20001102 (200107)
                                                C120001-00
ADT WO 2000063418 A1 WO 2000-US10173 20000414; AU 2000042455 A AU 2000-42455
    20000414
FDT AU 2000042455 A Based on WO 200063418
PRAI US 1999-292274 19990415
    ICM C12Q001-00
    ICS C07H019-20; C12Q001-02; C12Q001-04; C12Q001-32; G01N033-00;
    CPI EPI
CTCS CITATION COUNTERS
______
                      Cited Patents Count (by inventor)
PNC.DI
                      Cited Patents Count (by examiner)
PNC.DX
IAC.DI 0
                      Cited Issuing Authority Count (by inventor)
                      Cited Issuing Authority Count (by examiner)
IAC.DX 2
PNC.GI
                      Citing Patents Count (by inventor)
                      Citing Patents Count (by examiner)
PNC.GX
IAC.GI
                      Citing Issuing Authority Count (by inventor)
                      Citing Issuing Authority Count (by examiner)
IAC.GX
                      Cited Literature References Count (by inventor)
CRC.I
                      Cited Literature References Count (by examiner)
CRC.X
                     UPD: 20021122
CDP CITED PATENTS
    Cited by Examiner
    _____
    CITING PATENT CAT CITED PATENT ACCNO
    ______
    WO 200063418 A Y US 5656449 A 1997-414585/38
                  PA: (MOLE-N) MOLECULAR PROBES INC
                  IN: YUE, S T
                          US 5658751 A 1996-251457/25
                  PA: (MOLE-N) MOLECULAR PROBES INC
                  IN: HAUGLAND, R P; JIN, X; JONES, L J; MILLARD, P J;
                      MOZER, T J; POOT, M; ROTH, B L; SINGER, V L; YUE, S T;
                      POOT, M E
                          WO 9600902 A 1990-164056/21
                  PA: (SIHR-I) SHIRA K S
                  IN: SIHRA, K S
                          WO 9600902 A 1996-077582/08
                  PA: (BIOM-N) BIOMETRIC IMAGING INC; (LEEL-I) LEE L G;
                      (WOOS-I) WOO S L
                  IN: LEE, L G; WOO, S L
                          WO 9600902 Al 1996-077582/08
                      (BIOM-N) BIOMETRIC IMAGING INC; (LEEL-I) LEE L G;
                       (WOOS-I) WOO S L
                  IN: LEE, L G; WOO, S L
REN LITERATURE CITATIONS UPR: 20010221
    Citations by Examiner
     ______
```

CITED LITERATURE

CITING PATENT CAT

```
WO 200063418 A
                             CARDONE ET AL.: 'Phorbol myristate
                             acetate-mediated stimulation of transcytosis and
                             apical recycling in MDCK cells' THE JOURNAL OF
                             CELL BIOLOGY vol. 124, no. 5, March 1994, pages
                             717 - 727, XP002930076
                             LIPOWSKA ET AL.: 'New near-infrared cyanine dyes
     WO 200063418 A
                             for labelling of proteins' SYNTHETIC
                             COMMUNICATIONS vol. 23, no. 21, 1993, pages 3087 -
                             3094, XP002930077
    WO 200063418
                             DATABASE CAPLUS BIOMETRIC IMAGING INC. ACC. NO.
                             1996194739 LEE ET AL.: 'N-heteroaromatic ion and
                             iminium ion substituted cyanine dyes for use as
                             fluorescence labels' & WO 96 00902 A1
L113 ANSWER 3 OF 3 DPCI COPYRIGHT 2003 THOMSON DERWENT on STN
AN
     1998-042123 [04]
                        DPCI
DNC
    C1998-014108
     Ligand that binds the stalk of a cell's polymeric immunoglobulin receptor
TΙ
     - useful to target to, into or across mammalian epithelial cell
     biologically active component, e.g. nucleic acid, protein, lipid,
     carbohydrate, etc.
DC
     B04
    MOSTOV, K E; RICHMAN-EISENSTAT, J; MOSTOV, K
ΙN
     (REGC) UNIV CALIFORNIA
PΑ
CYC
    77
                  A1 19971211 (199804)* EN
                                              42p
                                                     C07K016-00
PΤ
    WO 9746588
                                                                       <--
        RW: AT BE CH DE DK EA ES FI FR GB GH GR IE IT KE LS LU MC MW NL OA PT
            SD SE SZ UG
         W: AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI GB GE
            HU IL IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG MK MN MW MX
            NO NZ PL PT RO RU SD SE SG SI SK TJ TM TR TT UA UG UZ VN YU
     AU 9730632
                   A 19980105 (199821)
                                                     C07K016-00
                                                                       <--
     EP 934338
                   A1 19990811 (199936) EN
                                                     C07K016-00
         R: AT BE CH DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE
                  A 19990630 (199944)
                                                     C07K016-00
     CN 1221428
                                                                       <--
                  A 20000328 (200023)
                                                                       <--
     US 6042833
                                                     A61K038-16
     JP 2000511432 W 20000905 (200047)
                                              46p
                                                     C12N015-09
                                                                       <--
                 B 20010111 (200108)
     AU 728587
                                                     C07K016-00
                                                                       <--
                  A 20010724 (200147)
                                                     C07K016-00
                                                                      <--
     IL 127238
     US 6340743
                  B1 20020122 (200208)
                                                     C07K016-28
                                                                      <--
                   C2 20021027 (200281)
                                                     C07K016-42
     RU 2191781
                                                                       <--
ADT WO 9746588 A1 WO 1997-US7944 19970514; AU 9730632 A AU 1997-30632
     19970514; EP 934338 A1 EP 1997-925515 19970514, WO 1997-US7944 19970514;
     CN 1221428 A CN 1997-195238 19970514; US 6042833 A Provisional US
     1996-18958P 19960604, US 1997-856383 19970514; JP 2000511432 W WO
     1997-US7944 19970514, JP 1998-500584 19970514; AU 728587 B AU 1997-30632
     19970514; IL 127238 A IL 1997-127238 19970514; US 6340743 B1 Provisional
     US 1996-18958P 19960604, Div ex US 1997-856383 19970514, US 1999-475088
     19991230; RU 2191781 C2 WO 1997-US7944 19970514, RU 1999-100279 19970514
FDT AU 9730632 A Based on WO 9746588; EP 934338 Al Based on WO 9746588; JP
     2000511432 W Based on WO 9746588; AU 728587 B Previous Publ. AU 9730632,
     Based on WO 9746588; US 6340743 B1 Div ex US 6042833; RU 2191781 C2 Based
     on WO 9746588
                      19960604; US 1997-856383 19970514; US 1999-475088
PRAI US 1996-18958P
     19991230
     ICM A61K038-16; C07K016-00; C07K016-28; C07K016-42; C12N015-09
IC
         A61K039-385; A61K039-395; C07K016-46; C12N015-13
     ICS
FS
     CPI
                                  UPE: 20020731
EXF EXAMINER'S FIELD OF SEARCH
```

NCL US 6042833 A 20000328

```
424/134.100; 424/185.100; 424/193.100; 530/380; 530/395; 530/403 US 6340743 B1 20020122 000/424.130 .1; 000/424.132 .1; 000/424.133 1-1351; 000/424.139 .1; 000/424.141 .1; 000/424.143 .1; 000/424.178 .1; 000/424.182 .1; 000/424.183 .1; 000/530.387 .1; 000/530.387 .3; 000/530.387 .5; 000/530.387 .9; 000/530.388 .1; 000/530.388 .22; 000/530.389 .1; 000/530.391 .1; 000/530.391 .3; 000/530.391 .7
```

CTCS CITATION COUNTERS

PNC.DI	0	Cited Patents Count (by inventor)
PNC.DX	0	Cited Patents Count (by examiner)
IAC.DI	0	Cited Issuing Authority Count (by inventor)
IAC.DX	0	Cited Issuing Authority Count (by examiner)
PNC.GI	0 .	Citing Patents Count (by inventor)
PNC.GX	2	Citing Patents Count (by examiner)
IAC.GI	0	Citing Issuing Authority Count (by inventor)
IAC.GX	2	Citing Issuing Authority Count (by examiner)
CRC.I	0	.Cited Literature References Count (by inventor)
CRC.X	34	Cited Literature References Count (by examiner)

CDP CITED PATENTS UPD: 20020731

Cited by Examiner

CITING PATENT	CAT	CITED PATENT	ACCNO
US 6042833 US 6340743	 А В1	No Citations No Citations	
WO 9746588	A	No Citations	

REN LITERATURE CITATIONS UPR: 20020731

Citations by Examiner

CITING PATENT	CAT ·	CITED LITERATURE
US 6042833 A		Mazanec et al., J. Virol. 69(2):1339-1343 (Feb. 1995).
US 6042833 A	•	Williams, G., Tibtech 6:36-42 (Feb. 1988).
US 6042833 A		Hudson, L. et al. (ed), Practical Immunology, 2nd edition, pp. 192-202, 1980.
US 6042833 A		Solari, R. et al., J. Biol. Chem. 260:1141-1145, Antibodies recognizing differen domains of the polymeric immunoglobulin receptor, 1985.
US 6042833 A	•	
US 6042833 A		Solari et al., J. Histochemistry and Cytochemistry 34(1):17-23 (1986).
US 6042833 A		Breitfeld et al., J. Cell Biology 109:475-486 (1989).
US 6042833 A		Piskurich et al., Journal of Immunology 154:1735-1747 (1995).
US 6042833 A		Ferkol et al., J. Clin. Invest. 95:493-502 (1995).
US 6042833 A		Wu et al., J. Biol. Chem. 262:4429-4432 (1987).
US 6042833 A		Breitfeld et al., Methods in Cell Biology 32:329-337 (1989).

US	6042833	A	Mostov et al., Ann. Rev. Immunol. 12:63-84 (1994).
	6042833	A	Ferkol et al., J. Clin. Invest. 92:2394-2400 (Nov.
0.5	0042033	А	
			1993).
US	6340743	B1	Piskurich et al., Journal of Immunology
			154:1735-1747 (1995).
US	6340743	B1	Breitfeld et al., J. Cell Biology 109:475-486
			(1989).
IIC	6340743	B1	Solari, R., et al., J. Biol. Chem. 260:1141-1145,
0.5	0540745	DI	Antibodies recognizing different domains of the
			polymeric immunoglobulin receptor. (1985).
US	6340743	B1	Mostov, Keith E., et al., Nature, 308(5954):37-43
			(Mar. 1, 1984).
US	6340743	B1	Mostov, Keith E., et al., Proc. Natl. Acad. Sci.,
		•	USA 77(12):7257-7261 (Dec. 1980).
IIC	6340743	В1	Solari et al., J. Histochemistry and Cytochemistry
0.5	0340743	DΙ	
			34(1):17-23 (1986).
US	6340743	B1	Ferkol et al., J. Clin. Invest. 92:2394-2400 (Nov.
			1993).
US	6340743	B1 '	Mostov et al., Ann. Rev. Immunol. 12:63-84 (1994).
US	6340743	B1	Ferkol et al., J. Clin. Invest. 95:493-502 (1995).
	6340743	B1	Wu et al., J. Biol. Chem. 262:4429-4432 (1987).
	6340743	B1	Breitfeld et al., Methods in Cell Biology
0.5	0340743	ĐI	
			32:329-337 (1989).
US	6340743	B1	Eiffert et al., Physiol. Chem. 365:1489-1495
			(1984).
US	6340743	В1	Mazanec et al., J. Virol. 69(2):1339-1343 (Feb.
			1005)
			1995).
· IIS	6340743	B1	1995). Williams, G., TIBTECH 6:36-42 (Feb. 1988).
	6340743 6340743	B1 B1	Williams, G., TIBTECH 6:36-42 (Feb. 1988).
	6340743 6340743	B1 B1	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd
US	6340743	B1	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980).
US			Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al.,
US	6340743	B1	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al., "Molecular Cloning of the Mouse Polymeric Ig
US	6340743	B1	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al., "Molecular Cloning of the Mouse Polymeric Ig Receptor", pages 1735-1747.
us wo	6340743	B1	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al., "Molecular Cloning of the Mouse Polymeric Ig
us wo	6340743 9746588	B1 A	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al., "Molecular Cloning of the Mouse Polymeric Ig Receptor", pages 1735-1747. J. VIROL., February 1995, Vol. 69, No. 2, MAZANEC
us wo	6340743 9746588	B1 A	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al., "Molecular Cloning of the Mouse Polymeric Ig Receptor", pages 1735-1747. J. VIROL., February 1995, Vol. 69, No. 2, MAZANEC et al., "Intracellular Neutralization of Influenza
us wo	6340743 9746588	B1 A	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al., "Molecular Cloning of the Mouse Polymeric Ig Receptor", pages 1735-1747. J. VIROL., February 1995, Vol. 69, No. 2, MAZANEC et al., "Intracellular Neutralization of Influenza Virus by Immunoglobulin A Anti-hemagglutinin
us Wo Wo	6340743 9746588 9746588	B1 A	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al., "Molecular Cloning of the Mouse Polymeric Ig Receptor", pages 1735-1747. J. VIROL., February 1995, Vol. 69, No. 2, MAZANEC et al., "Intracellular Neutralization of Influenza Virus by Immunoglobulin A Anti-hemagglutinin Monoclonal Antibodies", pages 1339-1343.
us Wo Wo	6340743 9746588	B1 A	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al., "Molecular Cloning of the Mouse Polymeric Ig Receptor", pages 1735-1747. J. VIROL., February 1995, Vol. 69, No. 2, MAZANEC et al., "Intracellular Neutralization of Influenza Virus by Immunoglobulin A Anti-hemagglutinin Monoclonal Antibodies", pages 1339-1343. TIBTECH, February 1988, Vol. 6, WILLIAMS G.,
us Wo Wo	6340743 9746588 9746588	B1 A	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al., "Molecular Cloning of the Mouse Polymeric Ig Receptor", pages 1735-1747. J. VIROL., February 1995, Vol. 69, No. 2, MAZANEC et al., "Intracellular Neutralization of Influenza Virus by Immunoglobulin A Anti-hemagglutinin Monoclonal Antibodies", pages 1339-1343. TIBTECH, February 1988, Vol. 6, WILLIAMS G., "Novel Antibody Reagents: Production and
US WO WO	6340743 9746588 9746588 9746588	B1 A	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al., "Molecular Cloning of the Mouse Polymeric Ig Receptor", pages 1735-1747. J. VIROL., February 1995, Vol. 69, No. 2, MAZANEC et al., "Intracellular Neutralization of Influenza Virus by Immunoglobulin A Anti-hemagglutinin Monoclonal Antibodies", pages 1339-1343. TIBTECH, February 1988, Vol. 6, WILLIAMS G., "Novel Antibody Reagents: Production and Potential", pages 36-42.
US WO WO	6340743 9746588 9746588	B1 A	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al., "Molecular Cloning of the Mouse Polymeric Ig Receptor", pages 1735-1747. J. VIROL., February 1995, Vol. 69, No. 2, MAZANEC et al., "Intracellular Neutralization of Influenza Virus by Immunoglobulin A Anti-hemagglutinin Monoclonal Antibodies", pages 1339-1343. TIBTECH, February 1988, Vol. 6, WILLIAMS G., "Novel Antibody Reagents: Production and Potential", pages 36-42. J. HISTOCHEMISTRY CYTOCHEMISTRY, 1986, Vol. 34,
US WO WO	6340743 9746588 9746588 9746588	B1 A A	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al., "Molecular Cloning of the Mouse Polymeric Ig Receptor", pages 1735-1747. J. VIROL., February 1995, Vol. 69, No. 2, MAZANEC et al., "Intracellular Neutralization of Influenza Virus by Immunoglobulin A Anti-hemagglutinin Monoclonal Antibodies", pages 1339-1343. TIBTECH, February 1988, Vol. 6, WILLIAMS G., "Novel Antibody Reagents: Production and Potential", pages 36-42. J. HISTOCHEMISTRY CYTOCHEMISTRY, 1986, Vol. 34,
US WO WO	6340743 9746588 9746588 9746588	B1 A A	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al., "Molecular Cloning of the Mouse Polymeric Ig Receptor", pages 1735-1747. J. VIROL., February 1995, Vol. 69, No. 2, MAZANEC et al., "Intracellular Neutralization of Influenza Virus by Immunoglobulin A Anti-hemagglutinin Monoclonal Antibodies", pages 1339-1343. TIBTECH, February 1988, Vol. 6, WILLIAMS G., "Novel Antibody Reagents: Production and Potential", pages 36-42. J. HISTOCHEMISTRY CYTOCHEMISTRY, 1986, Vol. 34, No. 1, SOLARI et al., "Distribution and Processing
US WO WO	6340743 9746588 9746588 9746588	B1 A A	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al., "Molecular Cloning of the Mouse Polymeric Ig Receptor", pages 1735-1747. J. VIROL., February 1995, Vol. 69, No. 2, MAZANEC et al., "Intracellular Neutralization of Influenza Virus by Immunoglobulin A Anti-hemagglutinin Monoclonal Antibodies", pages 1339-1343. TIBTECH, February 1988, Vol. 6, WILLIAMS G., "Novel Antibody Reagents: Production and Potential", pages 36-42. J. HISTOCHEMISTRY CYTOCHEMISTRY, 1986, Vol. 34, No. 1, SOLARI et al., "Distribution and Processing of the Polymeric Immunoglobulin Receptor in the
US WO WO	6340743 9746588 9746588 9746588	B1 A A	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al., "Molecular Cloning of the Mouse Polymeric Ig Receptor", pages 1735-1747. J. VIROL., February 1995, Vol. 69, No. 2, MAZANEC et al., "Intracellular Neutralization of Influenza Virus by Immunoglobulin A Anti-hemagglutinin Monoclonal Antibodies", pages 1339-1343. TIBTECH, February 1988, Vol. 6, WILLIAMS G., "Novel Antibody Reagents: Production and Potential", pages 36-42. J. HISTOCHEMISTRY CYTOCHEMISTRY, 1986, Vol. 34, No. 1, SOLARI et al., "Distribution and Processing of the Polymeric Immunoglobulin Receptor in the Rat Hepatocyte: Morphological and Biochemical
US WO WO	6340743 9746588 9746588 9746588	B1 A A	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al., "Molecular Cloning of the Mouse Polymeric Ig Receptor", pages 1735-1747. J. VIROL., February 1995, Vol. 69, No. 2, MAZANEC et al., "Intracellular Neutralization of Influenza Virus by Immunoglobulin A Anti-hemagglutinin Monoclonal Antibodies", pages 1339-1343. TIBTECH, February 1988, Vol. 6, WILLIAMS G., "Novel Antibody Reagents: Production and Potential", pages 36-42. J. HISTOCHEMISTRY CYTOCHEMISTRY, 1986, Vol. 34, No. 1, SOLARI et al., "Distribution and Processing of the Polymeric Immunoglobulin Receptor in the Rat Hepatocyte: Morphological and Biochemical Characterization of Subcellular Fractions", pages
wo wo wo	6340743 9746588 9746588 9746588	B1 A A A	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al., "Molecular Cloning of the Mouse Polymeric Ig Receptor", pages 1735-1747. J. VIROL., February 1995, Vol. 69, No. 2, MAZANEC et al., "Intracellular Neutralization of Influenza Virus by Immunoglobulin A Anti-hemagglutinin Monoclonal Antibodies", pages 1339-1343. TIBTECH, February 1988, Vol. 6, WILLIAMS G., "Novel Antibody Reagents: Production and Potential", pages 36-42. J. HISTOCHEMISTRY CYTOCHEMISTRY, 1986, Vol. 34, No. 1, SOLARI et al., "Distribution and Processing of the Polymeric Immunoglobulin Receptor in the Rat Hepatocyte: Morphological and Biochemical Characterization of Subcellular Fractions", pages 17-23.
wo wo wo	6340743 9746588 9746588 9746588	B1 A A	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al., "Molecular Cloning of the Mouse Polymeric Ig Receptor", pages 1735-1747. J. VIROL., February 1995, Vol. 69, No. 2, MAZANEC et al., "Intracellular Neutralization of Influenza Virus by Immunoglobulin A Anti-hemagglutinin Monoclonal Antibodies", pages 1339-1343. TIBTECH, February 1988, Vol. 6, WILLIAMS G., "Novel Antibody Reagents: Production and Potential", pages 36-42. J. HISTOCHEMISTRY CYTOCHEMISTRY, 1986, Vol. 34, No. 1, SOLARI et al., "Distribution and Processing of the Polymeric Immunoglobulin Receptor in the Rat Hepatocyte: Morphological and Biochemical Characterization of Subcellular Fractions", pages 17-23. J. CLIN. INVEST., November 1993, Vol. 92, FERKOL
wo wo wo	6340743 9746588 9746588 9746588	B1 A A A	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al., "Molecular Cloning of the Mouse Polymeric Ig Receptor", pages 1735-1747. J. VIROL., February 1995, Vol. 69, No. 2, MAZANEC et al., "Intracellular Neutralization of Influenza Virus by Immunoglobulin A Anti-hemagglutinin Monoclonal Antibodies", pages 1339-1343. TIBTECH, February 1988, Vol. 6, WILLIAMS G., "Novel Antibody Reagents: Production and Potential", pages 36-42. J. HISTOCHEMISTRY CYTOCHEMISTRY, 1986, Vol. 34, No. 1, SOLARI et al., "Distribution and Processing of the Polymeric Immunoglobulin Receptor in the Rat Hepatocyte: Morphological and Biochemical Characterization of Subcellular Fractions", pages 17-23. J. CLIN. INVEST., November 1993, Vol. 92, FERKOL et al., "Gene Transfer into Respiratory Epithelial
wo wo wo	6340743 9746588 9746588 9746588	B1 A A A	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al., "Molecular Cloning of the Mouse Polymeric Ig Receptor", pages 1735-1747. J. VIROL., February 1995, Vol. 69, No. 2, MAZANEC et al., "Intracellular Neutralization of Influenza Virus by Immunoglobulin A Anti-hemagglutinin Monoclonal Antibodies", pages 1339-1343. TIBTECH, February 1988, Vol. 6, WILLIAMS G., "Novel Antibody Reagents: Production and Potential", pages 36-42. J. HISTOCHEMISTRY CYTOCHEMISTRY, 1986, Vol. 34, No. 1, SOLARI et al., "Distribution and Processing of the Polymeric Immunoglobulin Receptor in the Rat Hepatocyte: Morphological and Biochemical Characterization of Subcellular Fractions", pages 17-23. J. CLIN. INVEST., November 1993, Vol. 92, FERKOL et al., "Gene Transfer into Respiratory Epithelial Cells by Targeting the Polymeric Immunoglobulin
wo wo wo	6340743 9746588 9746588 9746588	B1 A A A	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al., "Molecular Cloning of the Mouse Polymeric Ig Receptor", pages 1735-1747. J. VIROL., February 1995, Vol. 69, No. 2, MAZANEC et al., "Intracellular Neutralization of Influenza Virus by Immunoglobulin A Anti-hemagglutinin Monoclonal Antibodies", pages 1339-1343. TIBTECH, February 1988, Vol. 6, WILLIAMS G., "Novel Antibody Reagents: Production and Potential", pages 36-42. J. HISTOCHEMISTRY CYTOCHEMISTRY, 1986, Vol. 34, No. 1, SOLARI et al., "Distribution and Processing of the Polymeric Immunoglobulin Receptor in the Rat Hepatocyte: Morphological and Biochemical Characterization of Subcellular Fractions", pages 17-23. J. CLIN. INVEST., November 1993, Vol. 92, FERKOL et al., "Gene Transfer into Respiratory Epithelial Cells by Targeting the Polymeric Immunoglobulin Receptor", pages 2394-2400.
WO WO WO	6340743 9746588 9746588 9746588	B1 A A A	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al., "Molecular Cloning of the Mouse Polymeric Ig Receptor", pages 1735-1747. J. VIROL., February 1995, Vol. 69, No. 2, MAZANEC et al., "Intracellular Neutralization of Influenza Virus by Immunoglobulin A Anti-hemagglutinin Monoclonal Antibodies", pages 1339-1343. TIBTECH, February 1988, Vol. 6, WILLIAMS G., "Novel Antibody Reagents: Production and Potential", pages 36-42. J. HISTOCHEMISTRY CYTOCHEMISTRY, 1986, Vol. 34, No. 1, SOLARI et al., "Distribution and Processing of the Polymeric Immunoglobulin Receptor in the Rat Hepatocyte: Morphological and Biochemical Characterization of Subcellular Fractions", pages 17-23. J. CLIN. INVEST., November 1993, Vol. 92, FERKOL et al., "Gene Transfer into Respiratory Epithelial Cells by Targeting the Polymeric Immunoglobulin Receptor", pages 2394-2400.
WO WO WO	6340743 9746588 9746588 9746588 9746588	B1 A A A	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al., "Molecular Cloning of the Mouse Polymeric Ig Receptor", pages 1735-1747. J. VIROL., February 1995, Vol. 69, No. 2, MAZANEC et al., "Intracellular Neutralization of Influenza Virus by Immunoglobulin A Anti-hemagglutinin Monoclonal Antibodies", pages 1339-1343. TIBTECH, February 1988, Vol. 6, WILLIAMS G., "Novel Antibody Reagents: Production and Potential", pages 36-42. J. HISTOCHEMISTRY CYTOCHEMISTRY, 1986, Vol. 34, No. 1, SOLARI et al., "Distribution and Processing of the Polymeric Immunoglobulin Receptor in the Rat Hepatocyte: Morphological and Biochemical Characterization of Subcellular Fractions", pages 17-23. J. CLIN. INVEST., November 1993, Vol. 92, FERKOL et al., "Gene Transfer into Respiratory Epithelial Cells by Targeting the Polymeric Immunoglobulin Receptor", pages 2394-2400. ANN. REV. IMMUNOL., 1994, Vol. 12, MOSTOV et al.,
WO WO WO	6340743 9746588 9746588 9746588 9746588	B1 A A A	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al., "Molecular Cloning of the Mouse Polymeric Ig Receptor", pages 1735-1747. J. VIROL., February 1995, Vol. 69, No. 2, MAZANEC et al., "Intracellular Neutralization of Influenza Virus by Immunoglobulin A Anti-hemagglutinin Monoclonal Antibodies", pages 1339-1343. TIBTECH, February 1988, Vol. 6, WILLIAMS G., "Novel Antibody Reagents: Production and Potential", pages 36-42. J. HISTOCHEMISTRY CYTOCHEMISTRY, 1986, Vol. 34, No. 1, SOLARI et al., "Distribution and Processing of the Polymeric Immunoglobulin Receptor in the Rat Hepatocyte: Morphological and Biochemical Characterization of Subcellular Fractions", pages 17-23. J. CLIN. INVEST., November 1993, Vol. 92, FERKOL et al., "Gene Transfer into Respiratory Epithelial Cells by Targeting the Polymeric Immunoglobulin Receptor", pages 2394-2400. ANN. REV. IMMUNOL., 1994, Vol. 12, MOSTOV et al., "Transepithelial Transport of Immunoglobulins",
WO WO WO	6340743 9746588 9746588 9746588 9746588	B1 A A A	Williams, G., TIBTECH 6:36-42 (Feb. 1988). Hudson, L., et al. (ed), Practical Immunology, 2nd edition, pp. 192-202 (1980). J. IMMUNOL., 1995, Vol. 154, PISKURICH et al., "Molecular Cloning of the Mouse Polymeric Ig Receptor", pages 1735-1747. J. VIROL., February 1995, Vol. 69, No. 2, MAZANEC et al., "Intracellular Neutralization of Influenza Virus by Immunoglobulin A Anti-hemagglutinin Monoclonal Antibodies", pages 1339-1343. TIBTECH, February 1988, Vol. 6, WILLIAMS G., "Novel Antibody Reagents: Production and Potential", pages 36-42. J. HISTOCHEMISTRY CYTOCHEMISTRY, 1986, Vol. 34, No. 1, SOLARI et al., "Distribution and Processing of the Polymeric Immunoglobulin Receptor in the Rat Hepatocyte: Morphological and Biochemical Characterization of Subcellular Fractions", pages 17-23. J. CLIN. INVEST., November 1993, Vol. 92, FERKOL et al., "Gene Transfer into Respiratory Epithelial Cells by Targeting the Polymeric Immunoglobulin Receptor", pages 2394-2400. ANN. REV. IMMUNOL., 1994, Vol. 12, MOSTOV et al.,

CGP CITING PATENTS UPG: 20021009

Cited by Examiner

 	PATENT		 	 PATENT	ACCNO
		 А			2000-549134/52

PA: (DGIB-N) DGI BIOTECHNOLOGIES; (OKLA-N) OKLAHOMA MEDICAL RES FOUND; (TEXA) UNIV TEXAS SYSTEM; (DGIB-N)

DGI BIOTECHNOLOGIES INC

IN: CAPRA, J D; HEXHAM, J M; MANDECKI, W; WHITE, K

WO 9746588 A1 US 6207195 B1 1999-080847/01

PA: (UYJO) UNIV JOHNS HOPKINS SCHOOL MEDICINE; (UYJO) UNIV

JOHNS HOPKINS

IN: LEONG, K; RUBENSTEIN, R; WALSH, S; ZEITLIN, P;

LEONG, K W

=> fil hcaplus FILE 'HCAPLUS' ENTERED AT 08:09:40 ON 23 JUL 2003 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2003 AMERICAN CHEMICAL SOCIETY (ACS)

Copyright of the articles to which records in this database refer is held by the publishers listed in the PUBLISHER (PB) field (available for records published or updated in Chemical Abstracts after December 26, 1996), unless otherwise indicated in the original publications. The CA Lexicon is the copyrighted intellectual property of the the American Chemical Society and is provided to assist you in searching databases on STN. Any dissemination, distribution, copying, or storing of this information, without the prior written consent of CAS, is strictly prohibited.

FILE COVERS 1907 - 23 Jul 2003 VOL 139 ISS 4 FILE LAST UPDATED: 22 Jul 2003 (20030722/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> d bib abs tot 1145

L145 ANSWER 1 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN

AN 2001:519374 HCAPLUS

DN 135:121191

TI Bifunctional molecules for delivery of therapeutics

IN Davis, Pamela B.; Ferkol, Thomas W., Jr.; Eckman, Elizabeth

PA Case Western Reserve University, USA

SO U.S., 34 pp., Cont.-in-part of U.S. 6,072,041. CODEN: USXXAM

DT Patent

LA English

FAN.CNT 9

	PA	FENT	NO.		KII	ND	DATE			AF	PLI	CATI	ON NO	Э.	DATE			
ΡI	US	6261	787		B:	ŀ	2001	0717		US	19	99-2	6403	2	1999	0308		
	US	5972	900		Α		1999	1026		US	19	96-6	5570	5	1996	0603	<	
	US	5972	901		Α		1999	1026		US	19	96-6	5690	6	1996	0603		
	US	6072	041		Α		2000	0606		US	19	97-9	5733	3	1997	1024		
	WO	2000	0536	23	A.	1	2000	0914		WC	20	00-U	S593	0	2000	0308		
		W:	ΑU,	CA,	JP													
		RW:	AT,	BE,	CH,	CY,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,
			PT,	SE														
	ΕP	1165	597		A.	1	2002	0102		EF	20	00-9	1378	4	2000	0308		
		R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
			IE,															
	JΡ	2003	5221	17	T	2	2003	0722		JP	20	00-6	0405	8	2000	308		
PRAI	US	1996	-655	705	A.	2	1996	0603										

```
US 1996-656906
                      A2
                            19960603
                            19971024
     US 1997-957333
                      Α2
     US 1994-216534
                      В2
                            19940323
                      A1
     WO 1995-US3677
                            19950323
     US 1999-264032
                      Α
                            19990308
     WO 2000-US5930
                      W
                            20000308
     A bifunctional mol. consisting of a therapeutic mol. and a ligand
AΒ
     which specifically binds a transcytotic receptor can be transported
     specifically from the basolateral surface of epithelial cells to the
     apical surface. This approach provides the ability to deliver a
     therapeutic mol. directly to the apical surface of the epithelium, by
     targeting the transcytotic receptor with an appropriate ligand.
     Thus, the highest concn. of the therapeutic mol. will be at the apical
     surface, where it can have the greatest therapeutic effect.
              THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT 9
              ALL CITATIONS AVAILABLE IN THE RE FORMAT
L145 ANSWER 2 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN
     2000:573823 HCAPLUS
ΑN
     133:176176
· DN
     Polymeric immunoglobulin receptor (
TI
     pIgR) -binding domains and methods of use therefor
IN
     Capra, J. Donald; White, Kendra; Hexham, J. Mark; Mandecki, Wlodeck
     Oklahoma Medical Research Foundation, USA; Board of Regents, the
PΑ
     University of Texas System; Dgi Biotechnologies
     PCT Int. Appl., 139 pp.
SO
     CODEN: PIXXD2
DT
     Patent
T.A
     English
FAN.CNT 1
                     KIND DATE
                                          APPLICATION NO.
                                                           DATE
     PATENT NO.
                                          -----
                     ----
     _____
     WO 2000047611 A2
                                          WO 2000-US3650
                                                           20000211 <--
PΙ
                            20000817
                     A3
     WO 2000047611
                            20001130
            AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU,
             CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL,
             IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA,
             MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI,
             SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM,
             AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE,
             DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF,
             CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                                           20000211
     EP 1151000
                      A2
                           20011107
                                          EP 2000-906030
             AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO
                                           JP 2000-598527
                                                           20000211
     JP 2002539771
                      Т2
                            20021126
PRAI US 1999-119932P
                       P
                            19990212
     WO 2000-US3650
                      W
                            20000211
     The present invention identifies a domain located in the C.alpha.3 domain
ΑB
     of IgA that is responsible for targeting of the polymeric
     Ig receptor (pIgR) and transport of the
     antibody to the mucosal epithelium. This pIgR-binding domain
     may be used to target a wide variety of compns., including proteins,
     nucleic acids, drugs and diagnostic agents, to the mucosal surface.
     more specific targeting agents may be used in conjunction with the
     pIgR-binding domain to define further the ultimate localization of
     the complexes in the body. Treatment of a large no. of disease conditions
     such as viral, fungal and bacterial infections, as well as cancer, may be
     improved through the use of a pIgR-binding domain.
```

L145 ANSWER 3 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN AN 2000:381472 HCAPLUS

```
133:3719
DN
    Antibody fusion proteins for targeting apical epithelium
TI
    Davis, Pamela B.; Ferkol, Thomas; Eckman, Elizabeth; Schreiber, John; Luk,
IN
    John M.
    Case Western Reserve University, USA
PA
    U.S., 24 pp., Cont.-in-part of U.S. 655,705.
SO
    CODEN: USXXAM
DT
    Patent
    English
LA
FAN.CNT 9
                                                           DATE
    PATENT NO.
                     KIND DATE
                                          APPLICATION NO.
                                          _____
                                                           -----
    ______
                     ---- -----
                           20000606
                                          US 1997-957333
                                                           19971024
    US 6072041
                     Α
                                                           19960603 <--
    US 5972900
                     Α
                           19991026
                                          US 1996-655705
                                          US 1996-656906
                                                           19960603
    US 5972901
                     A
                           19991026
                                          US 1999-264032
    US 6261787
                     В1
                         20010717
                                                           19990308
                         20010911
    US 6287817
                     В1
                                          US 2000-559393
                                                           20000426
PRAI US 1996-655705
                     A2
                           19960603
    US 1996-656906
                      A2
                           19960603
    US 1994-216534
                      B2
                           19940323
    WO 1995-US3677
                      A1
                           19950323
    US 1997-957333
                      A2
                           19971024
ΑB
    The authors disclose the construction and characterization of single-chain
    antibody fusion proteins directed at the polymeric Ig
    receptor (pIgR). Such constructs have the ability to
    deliver a therapeutic protein directly to the apical surface of the
    epithelium. In one example, a fusion protein with .alpha.1-antitrypsin
    was transported to the apical surface of MDCK cells expressing a transgene
     for plgR.
             THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT 4
             ALL CITATIONS AVAILABLE IN THE RE FORMAT
L145 ANSWER 4 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN
ΑN
    1999:686690 HCAPLUS
DN
    131:327493
    Serpin enzyme complex receptor-mediated gene transfer
ΤI
    Ferkol, Thomas W., Jr.; Davis, Pamela B.; Ziady, Assem-galal
IN
    Case Western Reserve University, USA
PA
SO
    U.S., 81 pp., Cont.-in-part of U.S. Ser. No. 655,705.
    CODEN: USXXAM
DT
    Patent
    English
T.A
FAN.CNT 9
    PATENT NO.
                     KIND DATE
                                          APPLICATION NO. DATE
                     ____
                           -----
                                          -----
                                          US 1996-656906 19960603
                           19991026
    US 5972901
                      Α
ΡI
                                          WO 1995-US3677 19950323
     WO 9525809
                     A1
                           19950928
            AM, AT, AU, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE, ES, FI,
            GB, GE, HU, IS, JP, KE, KG, KP, KR, KZ, LK, LR, LT, LU, LV, MD,
            MG, MN, MW, MX, NL, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK,
            TJ, TT
         RW: KE, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT,
            LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE,
             SN, TD,
                    ΤG
     US 5972900
                           19991026
                                          US 1996-655705
                                                           19960603 <--
                                          WO 1997-US9858
     WO 9746100
                      A1
                           19971211
                                                           19970603
            AU, CA, JP
         RW: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
    AU 9733044
                      A1
                           19980105
                                          AU 1997-33044
                                                           19970603
    AU 720223
                           20000525
                      B2
                                                           19970603
     EP 1006797
                      A1
                           20000614
                                          EP 1997-928891
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
```

IE, FI

```
JP 1998-500875
                            20000919
                                                             19970603
     JP 2000512140
                       T2
     US 6072041
                            20000606
                                            US 1997-957333
                                                             19971024
                       Α
     US 6261787
                       В1
                            20010717
                                            US 1999-264032
                                                             19990308
     US 6287817
                       В1
                            20010911
                                            US 2000-559393
                                                             20000426
                       В2
                            19940323
PRAI US 1994-216534
                       Α1
                            19950323
     WO 1995-US3677
                       A2
     US 1996-655705
                            19960603
                            19960603
     US 1996-656906
                       Α
     WO 1997-US9858
                       W
                            19970603
                       A2
                            19971024
     US 1997-957333
```

Nucleic acids are compacted, substantially without aggregation, to AB facilitate their uptake by target cells of an organism to which the compacted material is administered. The nucleic acids may achieve a clin. effect as a result of gene expression, hybridization to endogenous nucleic acids whose expression is undesired, or site-specific integration so that a target gene is replaced, modified or deleted. The targeting may be enhanced by means of a target cell-binding moiety. The nucleic acid is preferably compacted to a condensed state.

THERE ARE 79 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 79 ALL CITATIONS AVAILABLE IN THE RE FORMAT

L145 ANSWER 5 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN

1999:547711 HCAPLUS

DN 131:285105

- In vitro transport of active .alpha.1-antitrypsin to the apical surface of TI epithelia by targeting the polymeric immunoglobulin
- ΑU Eckman, Elizabeth A.; Mallender, William D.; Szegletes, Tivadar; Silski, Catherine L.; Schreiber, John R.; Davis, Pamela B.; Ferkol, Thomas
- Department of Pediatrics, Case Western Reserve University, Cleveland, OH, CS 44106, USA
- SO American Journal of Respiratory Cell and Molecular Biology (1999), **21(2)**, **246**-252 CODEN: AJRBEL; ISSN: 1044-1549
- PB American Lung Association
- DΤ Journal
- LA English

AB

In cystic fibrosis (CF), the intense host inflammatory response to chronic infection largely accounts for the progressive pulmonary disease, and ultimately death. Neutrophils are the prominent inflammatory cells in the lungs of patients with CF, and large amts. of neutrophil elastase (NE) are released during phagocytosis. Besides having direct effects on structural elastin, NE stimulates the release of proinflammatory mediators from the respiratory epithelium and is a potent secretagogue. Therapeutic use of elastase inhibitors in CF has been complicated by difficulties in delivery to the crit. site in the airway-the surface of the epithelium. We describe a unique strategy to protect the respiratory epithelial cell surface directly by capitalizing on the nondegradative transcytotic pathway of the polymeric Ig receptor (pIgR). A recombinant fusion protein was constructed consisting of an antihuman pIgR single-chain Fv (scFv) antibody linked to human .alpha.1-antitrypsin (A1AT), an inhibitor of NE. The recombinant scFv-AlAT fusion protein bound specifically to the pIgR on the basolateral surface of an epithelial cell monolayer, and was transported and released into the apical medium where the AlAT domain was capable of forming an inactivation complex with NE. Thus, AlAT linked to an antihuman plgR scFv was delivered in receptor-specific fashion from the basolateral to apical surface and was released as an active antiprotease, indicating that it is feasible to deliver therapeutic proteins to the apical surface of epithelia by targeting the pIgR

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L145 ANSWER 6 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN

AN 1999:69867 HCAPLUS

DN 130:150635

TI Chemically reactive unsymmetrical cyanine dyes and their conjugates

IN Haugland, Richard P.; Singer, Victoria L.; Yue, Stephen T.; Millard, Paul J.

PA Molecular Probes, Inc., USA

SO U.S., 27 pp., Cont.-in-part of U.S. 5,658,751.

CODEN: USXXAM

DT Patent

LA English

FAN. CNT 8

FAN.	CNT 8				
	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
ΡI	US 5863753	A	19990126	US 1997-914439	19970819
	US 5658751	Α	19970819	US 1994-331031	19941027 <
PRAI	US 1994-331031	A2	19941027		
	US 1993-47683	B2	19930413		
	US 1994-90890	A2	19940712		
OS	MARPAT 130:15063	5			•
GI					

The invention comprises cyanine dyes, in particular chem. reactive dyes, conjugates of reactive cyanine dyes, the non-covalent complexes of nucleic acids with the dyes and dye-conjugates of the invention, and a method of forming a nucleic acid complex with the dyes and dye-conjugates of the present invention. The dyes of the invention are useful for the prepn. of dye-conjugates. The presence of a reactive group on the unsym. cyanine dyes of the invention facilitates their covalent conjugation to a variety of substances, both biol. and synthetic. Double-stranded DNA was photoaffinity labeled with I (prepn. given).

RE.CNT 28 THERE ARE 28 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L145 ANSWER 7 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN

AN 1999:7813 HCAPLUS

DN 130:71529

TI Therapeutic nanospheres containing sodium 4-phenylbutyrate for treatment of cystic fibrosis by CFTR gene therapy

IN Walsh, Scott; Rubenstein, Ronald; Zeitlin, Pamela; Leong, Kam

PA Johns Hopkins University School of Medicine, USA

SO PCT Int. Appl., 24 pp.

```
CODEN: PIXXD2
DT
    Patent
T.A
    English
FAN.CNT 1
                     KIND DATE
                                          APPLICATION NO.
                                                           DATE
                     ----
                                          _____
                     A2
                           19981217
                                         WO 1998-US11880 19980611
    WO 9856370
    WO 9956370
                     ΑЗ·
                           19990401
            AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE,
            DK, EE, ES, FI, GB, GE, GH, GM, GW, HU, ID, IL, IS, JP, KE, KG,
            KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX,
            NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT,
            UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
        RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES,
            FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI,
            CM, GA, GN, ML, MR, NE, SN, TD, TG
                                          CA 1998-2303268 19980611
                           19981217
    CA 2303268
                      AA
                                          AU 1998-80624
    AU 9880624
                      Α1
                           19981230
                                                           19980611
    AU 749032
                      В2
                           20020620
                                          EP 1998-928941
                                                           19980611
    EP 989849
                      A2
                           20000405
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
            IE, FI
                                          US 1998-95882
                                                           19980611 <--
    US 6207195
                           20010327
                                          JP 1999-503069
    JP 2002506436
                      T2
                           20020226
                                                           19980611
PRAI US 1997-49497P
                      Ρ
                           19970613
                     W
                           19980611
    WO 1998-US11880
    4-Phenylbutyrate exerts many beneficial biol. effects. It appears to
AΒ
    induce the transcription of certain promoters, as well as having a
    remedial effect on proteins which are aberrantly localized within the
    cell. In addn., it appears to cause cells to developmentally
    differentiate. The present invention provides nanosphere formulations of
     4-phenylbutyrate and other drugs which remediate defective protein
    localization intracellularly and can be used for treating cystic fibrosis.
     These formulations permit lower concns. of drugs to be administered,
    providing both cost and safety benefits.
L145 ANSWER 8 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN
    1997:574469 HCAPLUS
ΑN
DN
    127:231608
    Substituted unsymmetrical cyanine dyes with selected permeability
ΤI
    Yue, Stephen T.; Singer, Victoria L.; Roth, Bruce L.; Mozer, Thomas J.;
TN
    Millard, Paul J.; Jones, Laurie J.; Jin, Xiaokui; Haugland, Richard P.
    Molecular Probes, Inc., USA
PA
    U.S., 58 pp., Cont.-in-part of U.S. 5,436,134.
SO
    CODEN: USXXAM
DT
    Patent
    English
LΑ
FAN.CNT 8
     PATENT NO.
                     KIND DATE
                                          APPLICATION NO.
                                                           DATE
                     ----
                           -----
                                          ______
                                                           _____
     US 5658751
                            19970819
                                          US 1994-331031
                                                           19941027 <--
PΙ
     WO 9613552
                     A2
                           19960509
                                          WO 1995-US13706 19951027
         W: AU, CA, JP
         RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
    AU 9539672
                      A1
                            19960523
                                          AU 1995-39672
                                                           19951027
    AU 714890
                      B2
                            20000113
     WO 9613552
                      Α3
                            19960711
                                          WO 1995-EP13706 19951027
         W: AU, CA, JP
         RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
     EP 740689
                      A1
                          19961106
                                          EP 1995-937613
                                                           19951027
     EP 740689
                      В1
                            20020130
         R: AT, BE, CH, DE, FR, GB, LI, NL
                                          JP 1995-514689
    JP 09507879
                      T2
                          19970812
                                                           19951027
```

```
20020215
                                            AT 1995-937613
     AT 212653
                       Ε
                                                              19951027
                                            US 1997-914439
     US 5863753
                       Α
                             19990126
                                                              19970819
PRAI US 1993-47683
                       B2
                             19930413
     US 1994-90890
                       A2
                             19940712
     US 1994-331031
                       Α
                             19941027
                       W
                             19951027
     WO 1995-US13706
     MARPAT 127:231608
OS
```

The invention describes the prepn. and use of fluorescent stains for nucleic acids derived from unsym. cyanine dyes comprising a substituted benzazolium ring system linked by a methine bridge to a pyridinium or quinolinium ring system having at least one substituent on the pyridinium or quinolinium ring that contains a heteroatom. The presence of the heteroatom-contg. substituent results in higher sensitivity to oligonucleotides and larger nucleic acid polymers in a wide range of cells and gels, and for use in anal. of cell structure, membrane integrity or function. Thus, Dye 640 was prepd. by the methylation of 2-chloro-3-methylquinoline followed by the reaction of the intermediate iodide with 3-methyl-2-methylthiobenzothiazolinium tosylate in CH2C12 in the presence of 1 equiv. of NEt3. The use of these dyes in the detection of DNA in electrophoretic gels was demonstrated.

```
L145 ANSWER 9 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN
```

AN 1996:537669 HCAPLUS

DN 125:187585

TI Immunoglobulin fusion product with immunoglobulin receptor that protects Ig in mucosal environment, cDNA sequences, transgenic plants, and dental carie prevention

IN Hiatt, Andrew C.; Ma, Julian K.-C.; Lehner, Thomas

PA Planet Biotechnology, Inc., USA; United Medical and Dental Schools of Guy's and St. Thomas's Hospital

SO PCT Int. Appl., 154 pp. CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 2

```
KIND DATE
                                              APPLICATION NO.
                                                                 DATE
     PATENT NO.
                        ____
     WO 9621012
                              19960711
                                              WO 1995-US16889 19951227 <--
PΙ
                       A1
         W: AU, BR, CA, CN, CZ, FI, HU, JP, KR, MX, NO, NZ, PL, RU, SG RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
                              20000404
                                         . US 1995-434000
                                                                 19950504
     US 6046037
                       Α
     AU 9646088
                        A1
                              19960724
                                              AU 1996-46088
                                                                 19951227
                              20000810
     AU 722668
                        B2
                              19971119
                                               EP 1995-944237
                                                                 19951227
     EP 807173
                        Α1
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE
PRAI US 1994-367395
                      Α
                              19941230
     US 1995-434000
                        Α
                              19950504
     WO 1995-US16889
                        W
                              19951227
```

The Igs of the present invention are useful therapeutic Igs against mucosal pathogens such as S. mutans. The Igs contain a protection protein that protects the Igs in the mucosal environment. The invention also includes the greatly improved method of producing Igs in plants by producing the protection protein in the same cell as the other components of the Igs. The components of the Ig and assembled at a much improved efficiency. The method of the invention allows the assembly and high efficiency prodn. of such complex mols. The invention also contemplates the prodn. of Igs contg. protection proteins in a variety of cells, including plant cells, that can be selected for useful addnl. properties. The use of Igs contg. protection proteins as therapeutic antibodies against mucosal and other pathogens is also contemplated.

L145 ANSWER 10 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN AN 1996:443964 HCAPLUS

```
DN
     125:81256
     Substituted unsymmetrical cyanine dyes with selected permeability
TI
     Yue, Stephen T.; Singer, Victoria L.; Roth, Bruce L.; Mozer, Thomas J.;
IN
     Millard, Paul J.; Jones, Laurie J.; Jin, Xiaokui; Haugland, Richard P.;
     Poot, Martin
     Molecular Probes, Inc., USA
PA
     PCT Int. Appl., 85 pp.
SO
     CODEN: PIXXD2
DT
     Patent
     English
LA
FAN.CNT 8
                      KIND
                            DATE
                                           APPLICATION NO.
                                                             DATE
     PATENT NO.
                      ____
                            _____
                            19960509
                                           WO 1995-US13706
                                                             19951027
                       Α2
PΙ
     WO 9613552
         W: AU, CA, JP
         RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
                                           US 1994-331031
                                                             19941027 <--
     US 5658751
                       Α
                            19970819
                                           AU 1995-39672
                                                             19951027
                            19960523
     AU 9539672
                       Α1
                       B2
                            20000113
     AU 714890
                                           EP 1995-937613
                            19961106
                                                             19951027
     EP 740689
                       A1
     EP 740689
                       В1
                            20020130
         R: AT, BE, CH, DE, FR, GB, LI, NL
                                          JP 1995-514689
                                                             19951027
                       Т2
                            19970812
     JP 09507879
                                           AT 1995-937613
                                                             19951027
                       Ε
                            20020215
     AT 212653
PRAI US 1994-331031
                       Α
                            19941027
     US 1993-47683
                       B2
                            19930413
     US 1994-90890
                       Α2
                            19940712
                       W
                            19951027
     WO 1995-US13706
     MARPAT 125:81256
OS
     The invention describes the prepn. and use of fluorescent stains for
AB
     nucleic acids derived from unsym. cyanine dyes comprising a substituted
     benzazolium ring system linked by a methine bridge to a pyridinium or
     quinolinium ring system. The cyanine dyes of the invention possess a high
     sensitivity to oligonucleotides and larger nucleic acid polymers in a wide
     range of cells and gels, and are useful for the anal. of cell structure,
     membrane integrity or function, and detn. of cell cycle distribution.
L145 ANSWER 11 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN
     1996:194739 HCAPLUS
ΑN
     124:225822
DN
     N-heteroaromatic ion and iminium ion substituted
TΙ
     cyanine dyes for use as fluorescence labels
ΙN
     Lee, Linda G.; Woo, Sam L.
PΑ
     Biometric Imaging, Inc., USA
     PCT Int. Appl., 55 pp.
SO
     CODEN: PIXXD2
DT
     Patent
     English
LA
FAN.CNT 2
                                           APPLICATION NO.
     PATENT NO.
                      KIND DATE
                                                             DATE
                      ____
                            _----
                            19960111
                                           WO 1995-US8778
                                                            19950629 <--
     WO 9600902
                       Α1
PΙ
            AM, AT, AU, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE, ES, FI,
             GB, GE, HU, IS, JP, KE, KG, KP, KR, KZ, LK, LR, LT, LU, LV, MD,
             MG, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, TJ,
             TT, UA
         RW: KE, MW,
                    SD, SZ, UG, AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT,
             LU, MC,
                    NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE,
             SN, TD,
                     ТG
     US 5453505
                            19950926
                                            US 1994-268852
                                                             19940630
                       Α
     AU 9530085
                       Α1
                            19960125
                                            AU 1995-30085
                                                             19950629
                                                             19950629
     EP 769145
                       A1
                            19970423
                                           EP 1995-926272
         R: AT, BE, CH, DE, ES, FR, GB, IE, IT, LI, LU, NL, SE
```

```
PRAI US 1994-268852 19940630
US 1995-388607 19950214
WO 1995-US8778 19950629
```

OS MARPAT 124:225822

AΒ

GI For diagram(s), see printed CA Issue.

The present invention relates to iminium ion-substituted AΒ cyanine dyes having a fluorescence absorbance of between about 500 and 900 nm, a reduced tendency to aggregate and enhanced photostability. The cyanine dyes of the present invention are represented by formula I where n is 0, 1, 2 or 3; R1 and R2 are taken together to form an arom. ring or a fused polycyclic arom. ring; R3 and R4 are taken together to form an arom. ring or a fused polycyclic arom. ring; R5 and R6 are independently selected from the group consisting of (CH2)pX where p is 1-18 and X is a functional group that reacts with amino, hydroxy and sulfhydryl nucleophiles; R7 and R8 are independently selected from the group consisting of H, C1-C10 alkyl groups and where R7 and R8 are taken together to form a 5- or 6-membered heterocyclic ring; R9 are each independently selected from the group consisting of H, alkyl and where >1 R9 are taken together to form a 5- or 6-membered ring; Y is selected from the group consisting of C(CH3)2, S, O and Se; and Z is selected from the group consisting of C(CH3)2, S, O and Se. The present invention also relates to a method for using the cyanine dyes of the present invention to fluorescent label mols., particularly

```
biomols. such as antibodies, DNA, carbohydrates and cells.
L145 ANSWER 12 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN
ΑN
    1995:963703 HCAPLUS
DN
    123:332097
    Compacted nucleic acids and their delivery to cells for gene therapy
TI
    Hanson, Richard W.; Perales, Joseph C.; Ferkol, Thomas W., Jr.
IN
PA
    Ohio University, USA
SO
    PCT Int. Appl., 127 pp.
    CODEN: PIXXD2
DT
    Patent
LA
    English
FAN.CNT 9
    PATENT NO.
                     KIND DATE
                                          APPLICATION NO.
                                                          DATE
                     ____
     _____
                                          _____
    WO 9525809
                                    WO 1995-US3677 19950323
                    A1 19950928
ΡI
        W: AM, AT, AU, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE, ES, FI,
            GB, GE, HU, IS, JP, KE, KG, KP, KR, KZ, LK, LR, LT, LU, LV, MD,
            MG, MN, MW, MX, NL, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK,
            TJ, TT
         RW: KE, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT,
            LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE,
            SN, TD, TG
    CA 2186118
                           19950928
                                          CA 1995-2186118 19950323
                      AA \cdot
                                          AU 1995-21276
    AU 9521276
                      Α1
                           19951009
                                                           19950323
    AU 696455
                           19980910
                      В2
                                          EP 1995-914173
    EP 752005
                      A1
                           19970108
                                                           19950323
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, MC, NL, PT, SE
    JP 10503469
                     Т2
                           19980331
                                                          19950323
                                          JP 1995-524826
    US 5972900
                      Α
                           19991026
                                          US 1996-655705
                                                           19960603 <--
                    . A
    US 5972901
                                          US 1996-656906
                                                           19960603
                           19991026
    US 5877302
                      Α
                                          US 1997-716415
                                                           19970212
                           19990302
                      В1
                           20010313
                                          US 1998-217847
                                                           19981221
    US 6200801
PRAI US 1994-216534
                      Α
                           19940323
    WO 1995-US3677
                      W
                           19950323
    US 1996-655705
                      A2
                           19960603
    US 1996-655706
                      A2
                           19960603
                      A3
    US 1996-656096
                           19960603
```

Nucleic acids are compacted, substantially without aggregation, to facilitate their uptake by target cells of an organism to which the

compacted material is administered. The nucleic acids may achieve a clin. effect as a result of gene expression, hybridization to endogenous nucleic acids whose expression is undesired, or site-specific integration so that a target gene is replaced, modified or deleted. The targeting may be enhanced by means of a target-cell-binding moiety. The nucleic acid is preferably compacted to a condensed state.

```
L145 ANSWER 13 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN
    1995:875020 HCAPLUS
AN
    124:32008
DN
    N-Heteroaromatic ion- and iminium ion-substituted
TI ·
    cyanine dyes and their use as fluorescent labels
    Lee, Linda G.; Woo, Sam L.
ΙN
PΑ
    Biometric Imaging, Inc., USA
SO
    U.S., 18 pp.
    CODEN: USXXAM
DT
    Patent
LA
    English
FAN.CNT 2
                      KIND DATE
                                           APPLICATION NO.
    PATENT NO.
                                                            DATE
                                           -----
                                                            _____
                            _____
                            19950926
                                           US 1994-268852
                                                            19940630
PΙ
    US 5453505
                       Α
                                           WO 1995-US8778
                                                            19950629 <--
                            19960111
    WO 9600902
                      Α1
            AM, AT, AU, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE, ES, FI,
             GB, GE, HU, IS, JP, KE, KG, KP, KR, KZ, LK, LR, LT, LU, LV, MD,
             MG, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, TJ,
         RW: KE, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT,
             LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE,
             SN, TD, TG
                                           CA 1995-2194150 19950629
    CA 2194150
                            19960111
                            19960125
                                           AU 1995-30085
                                                            19950629
    AU 9530085
                       A1
    EP 769145
                       Α1
                            19970423
                                           EP 1995-926272
                                                            19950629
         R: AT, BE, CH, DE, ES, FR, GB, IE, IT, LI, LU, NL, SE
PRAI US 1994-268852
                            19940630
    US 1995-388607
                            19950214
    WO 1995-US8778
                            19950629
    MARPAT 124:32008
OS
    For diagram(s), see printed CA Issue.
GΙ
    The dyes having a fluorescence absorbance between 500 and 900.
AΒ
    nm, a reduced tendency to aggregate, and enhanced photostability.
    are represented by the formula I (A and B are arom. rings or fused
    polycyclic arom. rings; each R = H, alkyl, or 2 R together form a 5- or
     6-membered ring; R1, R2 = (CH2)pX; R3, R4 = H, C1-10 alkyl, or R3R4
    completes a 5- or 6-membered heterocyclic ring; X is a functional group
    that reacts with amino, OH, and SH nucleophiles; Z, Z1 = CMe2, S; m, n =
     0-3; p = 1-18). Thus, 2,3,3-trimethylindoline was alkylated with
    Br(CH2)5CO2H, condensed 2:1 with II, and the meso-Cl cyanine
     treated with 4-(dimethylamino) pyridine to give I [A = B = benzo, the R on
     the C atoms to either side of the meso C combine to form (CH2)3, the
     remaining R = H, R1 = R2 = (CH2)5CO2H, R3R4 = :CHC(NMe2):CH, Z = Z1 =
     CMe2, m = n = 1], absorption .lambda.max 786 nm, which was monoesterified
    with N-hydroxysuccinimide and used to label mouse IgG.
L145 ANSWER 14 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN
    1995:874985 HCAPLUS
ΑN
DN
     123:266187
    Method of encapsulating biological substances in microspheres
ΤI
     Tresco, Patrick A.; Mills, John F.
ΙN
     Brown University Research Foundation, USA
PA
SO
     U.S., 6 pp.
```

CODEN: USXXAM

Patent

DT

LA English FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE

PI US 5453368 A 19950926 US 1993-113778 19930827

US 5656469 A 19970812 US 1995-514780 19950814 <-
PRAI US 1993-113778 19930827

Amethod for encapsulating a biol. substance in biocompatible microcapsules, comprises (1) maintaining a coat-forming liq. film sheet comprising a polymer in an org. solvent, (2) causing droplets comprising biol. substance in an aq. medium to pass through the sheet to form microcapsules comprising cores of the droplets coated by the liq. film, and (3) permitting the microcapsules to pass through the sheet so that a portion of the polymer ppts. in the presence of water in the droplets while evapg. a portion of the solvent to form a continuous permeable polymer coating of sufficient structural integrity so that the microcapsules are self-supporting. A suitable app. is illustrated for performing the method of the present invention. Microencapsulation of PC 12 cells using polyacrylonitrile in DMF was demonstrated. A sample of the microcapsules was placed in culture and at the end of 6 wks, the microcapsules showed viable cells.

- L145 ANSWER 15 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN
- AN 1995:362852 HCAPLUS
- DN 122:158173
- TI Molecular cloning of the mouse **polymeric Ig**receptor. Functional regions of the molecule are conserved among
 five mammalian species
- AU Piskurich, Janet F.; Blanchard, May H.; Youngman, Kenneth R.; France, John A.; Kaetzel, Charlotte S.
- CS Inst. Pathol., Case Western Reserve Univ., Cleveland, OH, 44106, USA
- SO Journal of Immunology (1995), 154(4), 1735 -47
- CODEN: JOIMA3; ISSN: 0022-1767
 PB American Association of Immunologists
- DT Journal
- LA English
- AΒ Transcytosis of polymeric Ig (pIg) by mucosal epithelial cells is mediated by the polymeric Ig receptor (pIqR). Here the authors describe the characterization of a 3095-bp mouse plqR cDNA, which encodes a protein of 771 amino acids. Northern blot anal. detected a single mouse pIgR transcript of 3.9 kb, expressed at high levels in small intestine and liver, and at low levels in lung. Alignment of the amino acid sequences of mouse, rat, human, bovine, and rabbit pIgR revealed that functional regions of the mol. are conserved across species. In the extracellular region, conserved motifs include: a 23-amino acid pIg-binding site; 11 intradomain disulfide bonds, consensus sites for N-glycosylation, and a putative cleavage site at which the extracellular region of pIgR (secretory component) is released from the plasma membrane. A 10-amino acid sequence within the transmembrane region is highly conserved, possibly reflecting a mechanism for transmitting signals from the extracellular region to the cytoplasmic tail. Conversion within the cytoplasmic tail of pIqR is clustered in motifs that mediate polarized sorting, endocytosis, and transcytosis.
- L145 ANSWER 16 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN
- AN 1995:349785 HCAPLUS
- DN 122:124430
- TI Gene transfer into the airway epithelium of animals by targeting the polymeric immunoglobulin receptor
- AU Ferkol, Thomas; Perales, Jose C.; Eckman, Elizabeth; Kaetzel, Charlotte S.; Hanson, Richard W.; Davis, Pamela B.

CS Dep. Pediatr., Rainbow Babies Child. Hosp., Cleveland, OH, 44106, USA Journal of Clinical Investigation (1995), 95(2), 493-502

CODEN: JCINAO; ISSN: 0021-9738

- PB Rockefeller University Press
- DT Journal
- LA English
- AB Genes of interest can be targeted specifically to respiratory epithelial cells in intact animals with high efficiency by exploiting the receptor-mediated endocytosis of the polymerin

Ig receptor. A DNA carrier, consisting of the Fab portion of polyclonal antibodies raised against rat secretory component covalently linked to poly-L-lysine, was used to introduce plasmids contg. different reporter genes into airway epithelial cells in vivo. We obsd. significant levels of luciferase enzyme activity in protein exts. from the liver and lung, achieving max. values of 13,795 .+-. 4,431 and 346,954 .+-. 199,120 integrated light units (ILU) per mg of protein ext., resp. No luciferase activity was detected in spleen or heart, which do not express the receptor. Transfections using complexes consisting of an irrelevant plasmid (pCMV lacZ) bound to the bona fide carrier based on an irrelevant Fab fragment tissues resulted in background levels of luciferase activity in all tissues examd. Thus, only tissues that contain cells bearing the polymeric Ig receptor are

transfected, and transfection cannot be attributed to the nonspecific uptake of an irrelevant carrier-DNA complex. Specific mRNA from the luciferase gene was also detected in the lungs of transfected animals. To det. which cells in the lung are transfected by this method, DNA complexes were prepd. contg. expression plasmids with genes encoding the bacterial .beta.-galactosidase or the human interleukin 2 receptor.

Expression of these genes were localized to the surface epithelium of the airways with submucosal glands, and not the bronchioles and alveoli. Receptor-mediated endocytosis can be used to introduce functional genes into the respiratory epithelium of rats, and may be a useful technique for gene therapy targeting the lung.

- L145 ANSWER 17 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN
- AN 1995:335992 HCAPLUS
- DN 122:103619
- TI Intracellular neutralization of influenza virus by immunoglobulin A anti-hemagglutinin monoclonal antibodies
- AU Mazanec, Mary B.; Coudret, Christina L.; Fletcher, David R.
- CS Dep. Med. Pathol., Case Western Reserve Univ., Cleveland, OH, 44106, USA
- SO Journal of Virology (1995), 69(2), 1339-43
 - CODEN: JOVIAM; ISSN: 0022-538X
 B American Society for Microbiology
- PB America: DT Journal
- LA English
- Traditionally, IgA was thought to neutralize virus by forming complexes AB with viral attachment proteins, blocking attachment of virions to host epithelial cells. Recently we have proposed an intracellular action for dimeric IgA, which is actively transported through epithelial cells by the polymeric Ig receptor (pIgR), in that it may be able to bind to newly synthesized viral proteins within the cell, preventing viral assembly. To this effect, we have previously demonstrated that IqA monoclonal antibodies against Sendai virus, a parainfluenza virus, colocalize with the viral hemagglutinin-neuraminidase protein within infected epithelial cells and reduce intracellular viral titers. Here we det. whether IgA can interact with influenza virus hemagglutinin (HA) protein within epithelial cells. Polarized monolayers of Madin-Darby canine kidney epithelial cells expressing the plgR were infected on their apical surfaces with influenza virus A/Puerto Rico/8-Mount Sinai. Polymeric IgA anti-HA, but not IgG anti-HA,

delivered to the basolateral surface colocalized with HA protein within the cell by immunofluorescence. Compared with those of controls, viral titers were reduced in the supernatants and cell lysates from monolayers treated with anti-HA IgA but not with anti-HA IgG. Furthermore, the addn. of anti-IgA antibodies to supernatants did not interfere with the neutralizing activity of IgA placed in the basal chamber, indicating that IgA was acting within the cell and not in the extracellular medium to interrupt viral replication. Thus, these studies provide addnl. support for the concept that IgA can inhibit replication of microbial pathogens intracellularly.

```
L145 ANSWER 18 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN
```

AN 1994:296085 HCAPLUS

DN 120:296085

TI Transepithelial transport of immunoglobulins

AU Mostov, Keith E.

CS Dep. Anat., Univ. California, San Francisco, CA, 94143-0452, USA

SO Annual Review of Immunology (1994), 12, 63
-84

CODEN: ARIMDU; ISSN: 0732-0582

DT Journal; General Review

LA English

AB A review with 90 refs. Igs are transported across a variety of epithelial tissues. The best studied example of this is the transport of polymeric IgA and IgM by the polymeric Ig receptor (pIgR) across many types of epithelial cells. Transcytosis may be regulated by the heterotrimeric Gs protein, protein kinase C and calmodulin. IgG is transcytosed from the apical to basolateral surface in several epithelial tissues such as the placenta and the small intestine of newborn rats. The receptor for intestinal transport of IgG is structurally similar to class I MHC mols.

L145 ANSWER 19 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN

AN 1994:130640 HCAPLUS

DN 120:130640

TI Phorbol myristate acetate-mediated stimulation of transcytosis and apical recycling in MDCK cells

AU Cardone, Michael H.; Smith, Bradley L.; Song, Wenxia; Mochley-Rosen, Daria; Mostov, Keith E.

CS Dep. Anat., Univ. California, San Francisco, CA, 94143-0452, USA

SO Journal of Cell Biology (1994), 124(5), 717-28
CODEN: JCLBA3; ISSN: 0021-9525

DT Journal

LA English

Phorbol myristate acetate (PMA) stimulates transcytosis of the AB polymeric Ig receptor (pIgR) in MDCK cells. Apical release of pre-endocytosed ligand (dimeric IgA) bound to the pIgR can be stimulated 2-fold within 7 min of addn. of PMA while recycling of the ligand from the basal surface is not affected. In addn., apical surface delivery of plgR and cleavage of its ectodomain to secretory component (SC) is also stimulated by PMA. The recycling of apically internalized ligand back to the apical surface is similarly stimulated. These results suggest that the stimulation of apical delivery is from an apical recycling compartment. The effect of PMA suggests that protein kinase C (PKC) is involved in the regulation of pIgR trafficking in MDCK cells. To test this the authors down regulated PKC activity by pre-treating cells with PMA for 16 h and obsd. that transcytosis could no longer be stimulated by PMA. Western blots show that the PKC isoenzymes .alpha. and to a lesser extent .epsilon., are depleted from MDCK cells which have been pre-treated with PMA for 16 h and that treatment of MDCK cells with PMA for 5 min causes a dramatic translocation of the PKC .alpha. isoenzyme and

a partial translocation of the .epsilon. isoenzyme from the cytosol to the membrane fraction of cell homogenates. This translocation suggests that the .alpha. and/or .epsilon. isoenzymes may be involved in PMA-mediated stimulation of transcytosis. A mutant pIgR in which serines 664 and 726, the major sites of phosphorylation, are replaced by alanine is stimulated to transcytose by PMA, suggesting that phosphorylation of pIgR at these sites is not required for the effect of PMA. These results suggest that PMA-mediated stimulation of pIgR transcytosis may involve the activation of PKC .alpha. and/or .epsilon., and that this stimulation occurs independently of the major phosphorylation sites on the pIgR. Finally, PMA stimulates transcytosis of basolaterally internalized transferrin, suggesting that PMA acts to generally stimulate delivery of endocytosed proteins to the apical surface.

L145 ANSWER 20 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN

AN 1994:56667 HCAPLUS

DN 120:56667

TI New near-infrared cyanine dyes for labeling of proteins

AU Lipowska, Malgorzata; Patonay, Gabor; Strekowski, Lucjan

CS Dep. Chem., Georgia State Univ., Atlanta, GA, 30303, USA

SO Synthetic Communications (1993), 23(21),

3087-94

CODEN: SYNCAV; ISSN: 0039-7911

DT Journal

LA English

GΙ

AB Isothiocyanato-functionalized cyanine dyes I (X = 0, S) for labeling of proteins at amino groups are synthesized. The dyes and their adducts with amines show strong absorbance and fluorescence in the near-IR region of 750-850 nm.

Ι

L145 ANSWER 21 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN

AN 1994:1713 HCAPLUS

DN 120:1713

TI Gene transfer into respiratory epithelial cells by targeting the polymeric immunoglobulin receptor

AU Ferkol, Thomas; Kaetzel, Charlotte S.; Davis, Pamela B.

CS Dep. Pediatr., Rainbow Babies Child. Hosp., Cleveland, OH, 44106, USA

SO Journal of Clinical Investigation (1993), 92(5), 2394-400

CODEN: JCINAO; ISSN: 0021-9738

DT Journal

LA English

AB A system for targeting foreign DNA to epithelial cells in vitro has been developed by exploiting receptor-mediated endocytosis. The

polymeric Iq receptor transports dimeric IqA and IgM through epithelial cells, including those of the respiratory tract, by binding the Iqs at the basolateral surface and transporting them across the cell. Fab fragments of antibodies directed against the extracellular portion of the receptor, secretory component, are similarly transported. Anti-human secretory component Fab fragments were covalently linked to a polycation, and complexed to various expression plasmids. When bound to an expression plasmid contg. the Escherichia coli lacZ gene ligated to the Rous sarcoma virus promoter, the complexes transfected HT29.74 human colon carcinoma cells induced to express polymeric Ig receptor, but not those lacking the receptor. Primary cultures of human tracheal epithelial cells grown on collagen gels, which induce the expression of polymeric Ig receptor, were also transfected with the complexes. From 5 to 66% of the respiratory epithelial cells had .beta.-galactosidase activity after treatment, comparable to the percentage of cultured human tracheal epithelial cells that express polymeric Ig receptor (8-35%). The addn. of excess human secretory component (Fab ligand) to the culture medium at the time of transfection blocked the delivery of DNA. expression plasmid, either alone, complexed to the polycation, or complexed to a carrier based on an irrelevant Fab fragment, was not effective in transfecting either cell type. This DNA carrier system introduces DNA specifically into epithelial cells that contain pIqR in vitro.

L145 ANSWER 22 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN

AN 1993:464318 HCAPLUS

DN 119:64318

- TI Molecular cloning and exon-intron mapping of the gene encoding human transmembrane secretory component (the poly-Ig receptor)
- AU Krajci, Peter; Kvale, Dag; Tasken, Kjetil; Brandtzaeg, Per
- CS Lab. Immunohistochem., Norway
- SO European Journal of Immunology (1992), 22(9), 2309-15 CODEN: EJIMAF; ISSN: 0014-2980
- DT Journal
- LA English
- Secretory component (SC or the poly-Ig receptor) plays ΑB a crucial role in mucosal immunity by translocating polymeric IgA and IgM through secretory epithelial cells into external body fluids. Labeled restriction fragments from human SC cDNA were used to screen a human genomic leukocyte library. Three overlapping clones, spanning a total of 19 kb of the human SC gene, including 3 kb of the 5' flanking region, were characterized. The putative TATA box candidate, preceded by a CAAT-like box, was found 329 nucleotides upstream of the first exon. Altogether 11 exons covering the entire coding region were identified. The exon size ranged from 59 to 657 nucleotides and exon-intron junctions followed known consensus sequences. Three of the five extracellular Iq-related domains (D1, D4 and D5) were confined to one exon each (E3, E5 and E6), whereas D2 and D3 were encoded by the same exon (E4). The latter exon corresponds to that involved in alternate splicing of The membrane-spanning segment was confined to part of one exon rabbit SC. (E8). The cytoplasmic tail was encoded by four exons (E8-E11), whose boundaries encompassed fairly well the structural determinants proposed to be responsible for intracellular sorting of SC in the rabbit. The polymorphic restriction site reported earlier for PvuII was localized to the third intron.
- L145 ANSWER 23 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN
- AN 1990:455160 HCAPLUS
- DN 113:55160
- TI Expression and analysis of the polymeric immunoglobulin

receptor in Madin-Darby canine kidney cells using retroviral
vectors

- AU Breitfeld, Philip P.; Casanova, James E.; Harris, Jeanne M.; Simister, Neil E.; Mostov, Keith E.
- CS Med. Sch., Univ. Massachusetts, Worcester, MA, 01655, USA
- SO Methods in Cell Biology (1989), 32 (Vesicular Transp., Pt. B), 329-37 CODEN: MCBLAG; ISSN: 0091-679X
- DT Journal; General Review
- LA English
- AB A review with 10 refs. describes method for studying the expression and transport of the polymeric Ig receptor (poly-IgR) in Madin-Darby canine kidney (MDCK) cells. Topic covered were expression of the Poly-IgR in MDCK cells, prodn. of antibody against rabbit secretory component, labeling of cells producing Poly-IgR and immunopptn., growth of cells on filters, pulse-chase anal. of cells on filters, and measurement of transcytosis.
- L145 ANSWER 24 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN
- AN 1989:495175 HCAPLUS
- DN 111:95175
- TI Postendocytotic sorting of the ligand for the polymeric immunoglobulin receptor in Madin-Darby canine kidney cells
- AU Breitfeld, Philip P.; Harris, Jeanne M.; Mostov, Keith E.
- CS Whitehead Inst. Biomed. Res., Cambridge, MA, 02142, USA
- SO Journal of Cell Biology (1989), 109(2), 475 -86
- CODEN: JCLBA3; ISSN: 0021-9525 DT Journal
- LA English
- AΒ The polymeric Ig receptor (pIg-R) is responsible for the receptor-mediated transcytosis of polymeric Igs (IgA and IgM) across various epithelia. The present study investigated the postendocytotic pathway of the ligand for the pIg-R. After a 5-min internalization at the basolateral surface, .apprx.45% of internalized ligand recycles to the basolateral medium and 30% is transcytosed to the apical medium. Why transcytosis of ligand is unidirectional, going only from basolateral to apical, but not from apical to basolateral, was also examd. Several factors could explain this, such as proteolytic cleavage of the pIq-R at the apical surface, decreased apical endocytosis of ligand, or an intracellular sorting event. The protease inhibitor, leupeptin, inhibits the cleavage of the pIg-R but does not alter the unidirectionality of transcytosis. In addn., there is a significant amt. of apical endocytosis of ligand (70% of that obsd. basolaterally). Apically endocytosed ligand can return only to the apical surface. Thus, once ligand reaches the apical surface, it is trapped and cannot return to the basolateral It is proposed that unidirectionality of transcytosis is the result of intracellular sorting, and that this results from a signal(s) present on the pIg-R.
- L145 ANSWER 25 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN
- AN 1988:184690 HCAPLUS
- DN 108:184690
- TI Novel antibody reagents: production and potential
- AU Williams, Gareth
- CS MRC Lab. Mol. Biol., Univ. Postgrad. Med. Sch., Cambridge, CB2 2QH, UK
- SO Trends in Biotechnology (1988), 6(2),
 - CODEN: TRBIDM; ISSN: 0167-7799

- DT Journal; General Review
- LA English
- AB A review with 40 refs. By use of genetic engineering and special hybridomas, monoclonal antibodies with dual specificities, predetd. specificities, or addnl. functional moieties can be produced.
- L145 ANSWER 26 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN
- AN 1987:170223 HCAPLUS
- DN 106:170223
- TI Receptor-mediated in vitro gene transformation by a soluble DNA carrier system
- AU Wu, George Y.; Wu, Catherine H.
- CS Sch. Med., Univ. Connecticut, Farmington, CT, 06032, USA
- SO Journal of Biological Chemistry (1987), 262(10),

CODEN: JBCHA3; ISSN: 0021-9258

- DT Journal
- LA English
- Foreign DNA can be specifically delivered to cells by a sol. carrier AB system that takes advantage of receptor-mediated endocytosis. The expts. were based on the following concepts: (1) hepatocytes possess a unique receptor that binds and internalizes galactose-terminal (asialo-)glycoproteins; (2) DNA can bind to polycations in a strong but noncovalent manner forming sol. complexes; and (3) the gene for chloramphenicol acetyltransferase, a bacterial enzyme that acetylates chloramphenicol, is not present in mammalian cells. Asialoorosomucoid (ASOR) was coupled to poly-L-lysine to form an asialoorosomucoid-poly-Llysine conjugate. The plasmid, pSV2 CAT, was complexed to the conjugate in a molar ratio of 1:2. To test this complex, a model system was used consisting of hepatoma cell lines, Hep G2, asialoglycoprotein receptor (+), and SK-Hep 1, receptor (-). Each cell line was incubated with filtered ASOR.cntdot.poly-L-lysine.cntdot.DNA complex, or controls consisting of DNA plus ASOR, DNA plus poly-L-lysine, or DNA alone. were assayed for the presence of chloramphenicol acetyltransferase activity as a measure of gene transformation. SK-Hep 1, receptor (-) cells, produced no detectable acetylated chloramphenicol derivs. under any condition. However, Hep G2, receptor (+) cells, incubated with the ASOR.cntdot.poly-L-lysine.cntdot.DNA complex were transformed as indicated by the presence of chloramphenical acetyltransferase activity (0.028 chloramphenicol acetyltransferase units/106 cells). Mixts. of individual components of the complex failed to transform these cells. Competition by a 10-fold excess of ASOR prevented gene transformation by the ASOR.cntdot.poly-L-lysine.cntdot.DNA complex.
- L145 ANSWER 27 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN
- AN 1986:127893 HCAPLUS
- DN 104:127893
- TI Distribution and processing of the **polymeric**immunoglobulin receptor in the rat hepatocyte:
 morphological and biochemical characterization of subcellular fractions
- AU Solari, Roberto; Racine, Liliane; Tallichet, Corinne; Kraehenbuhl, Jean Pierre
- CS Swiss Inst. Exp. Cancer Res., Epalinges, 1066, Switz.
- SO Journal of Histochemistry and Cytochemistry (1986), 34 (1), 17-23 CODEN: JHCYAS; ISSN: 0022-1554
- DT Journal
- LA English
- AB Rat liver microsomes were fractionated and analyzed by immunochem. techniques for the IgA receptor (secretory component transmembrane form). The fraction enriched in the plasma membrane and rough endoplasmic reticulum contained predominantly a low-mol.-wt. form of the receptor [105 kilodaltons (kd)] which represents a core-glycosylated intermediate. In

the Golgi-enriched fraction, the receptor is present in its terminally glycosylated form and appears as a doublet with a mol. wt: of 115 kd. A lysosome-rich fraction contains both the 115 kd receptor and a 34 kd protein that was demonstrated by peptide mapping to be the membrane-anchoring domain of the receptor. Bile contains 31 kd and 29 kd proteins and hepatocyte cytosol contains a 32 kd protein that reacts with receptor-specific monoclonal antibody.

- L145 ANSWER 28 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN
- AN 1985:94102 HCAPLUS
- DN 102:94102
- TI Antibodies recognizing different domains of the polymeric immunoglobulin receptor
- AU Solari, Roberto; Kuehn, Lukas; Kraehenbuhl, Jean Pierre
- CS Inst. Biochim., Univ. Lausanne, Epalinges, CH-1066, Switz.
- SO Journal of Biological Chemistry (1985), 260(2), 1141-5
 CODEN: JBCHA3; ISSN: 0021-9258
- DT Journal
- LA English
- The receptor responsible for the transepithelial transport of IgA dimer AB antibodies is a transmembrane glycoprotein known as membrane secretory component (SCm). During transport, the membrane anchoring domain is cleaved and the ectoplasmic domain of the receptor (SCs) remains tightly bound to the IgA dimer in exosecretions. Monoclonal antibodies were produced with distinct specificities against both cytoplasmic and ectoplasmic epitopes of rabbit liver SCm. One antibody (anti-SC303) reacted both with SCm and free SCs but not with SCs bound to IgA dimer Therefore, it recognized an epitope close to the IgA dimer binding site. The other monoclonal antibody (anti-SC166), which was unable to react with SCs, bound to the 15-kilodalton cytoplasmic extension of the membrane-spanning domain of the receptor. A polyclonal antibody (GaR-SC), raised in a goat against rabbit milk SCs, reacted with a subpopulation of SCs not recognized by the anti-SC303 monoclonal antibody and in addn. also reacted with covalently bound sIgA. The 3 antibodies cross-reacted with rat SCm. The ability of the anti-SC166 monoclonal antibody to immunoadsorb subcellular organelles as a result of the cytoplasmic orientation of its epitope thus is demonstrated. These data indicate that there are functional differences between the high- and low-mol.-wt. families of SC in terms of IgA dimer binding.
- L145 ANSWER 29 OF 31 HCAPLUS COPYRIGHT 2003 ACS on STN .
- AN 1985:94074 HCAPLUS
- DN 102:94074
- TI The primary structure of the human free secretory component and the arrangement of the disulfide bonds
- AU Eiffert, Helmut; Quentin, Elmar; Decker, Joachim; Hillemeir, Sabine; Hufschmidt, Margarethe; Klingmueller, Dietrich; Weber, Michael H.; Hilschmann, Norbert
- CS Abt. Immunchem., Max-Planck-Inst. Exp. Med., Goettingen, D-3400, Fed. Rep. Ger.
- SO Hoppe-Seyler's Zeitschrift fuer Physiologische Chemie (1984), 365(12), 1489-95 CODEN: HSZPAZ; ISSN: 0018-4888
- DT Journal
- LA German
- The amino-acid sequence and the arrangement of the disulfide bonds of the free secretory component, isolated from colostrum from different women, were completely elucidated by the methods of protein chem. The free secretory component is a monomeric glycoprotein (mol. wt. .apprx.86,000), consisting of 558 amino acids with 7 carbohydrate chains bound to asparagine. The protein contains 20 cysteine residues but, as a special feature, no methionine. The polypeptide chain is divided into 5 regions

```
E MOSTOV K/AU
            121 S E3-E8
L2
                E CHAPIN S/AU
             23 S E6,E10-E12
L3
                E RICHMAN EISENSTAT J/AU
              8 S E3-E6
L4
             18 S E21, E22, E30, E32
L5
                E EISENSTAT/AU
                E LIGAND/CT
                E E38+ALL
L6
          15394 S E1
         330572 S LIGAND
L7
                E IMMUNOGLOBULIN RECEPTOR/CT
            185 S E89, E90
                E E4+ALL
           1556 S E10-E12
L9
L10
             62 S E85
L11
            386 S E9 (L) POLYM?
                E PIGR
            206 S E3
L12
L13
           1316 S IMMUNOGLOB? (L) RECEPTOR (L) ?POLYM?
L14
            366 S L6, L7 AND L8-L13
                E IMMUNOGLOBULINS/CT
           6667 S E3 (L) FRAGMENT?
L15
             25 S L14 AND L15
L17
            366 S (L6 OR L7 OR ?LIGAND?) AND L8-L13
L18
             25 S L15 AND L17
                E ANIMAL CELL/CT
          36281 S E3
L19
                E ANIMAL ORGAN/CT
                E E3+ALL
                E E2+ALL
L20
          30673 S E4,E5,E3
              9 S L17 AND L19
L21
              1 S L17 AND L20
L22
              9 S L21, L22
L23
               6 S L18 AND L23
L24
                E ANTIBOD/CT
                E E58+ALL
L25
         154078 S ANTIBODIES/CT
L26
            330 S L25 AND L8-L13
L27
             44 S L26 AND L15
L28
              5 S L27 AND L19, L20
L29
              7 S L24, L28
             39 S L2-L5 AND (L6 OR L7 OR ?LIGAND? OR L25 OR ANTIBOD?)
L30
L31
             26 S L30 AND L8-L13
L32
              3 S L30 AND L15
L33
             27 S L31, L32
L34
              2 S L33 AND L19, L20
L35
              7 S L29, L34
             25 S L33 NOT L35
L36
L37
             36 S L2-L5 AND ?PIGR?
L38
             49 $ L33-L37
L39
             12 S L30 NOT L38
             12 S L2 AND L3-L5
              2 S L3 AND L4, L5
L42
             12 S L40, L41
L43
              4 S L42 AND L38
             45 S L38 NOT L43
L44
             39 S L44 AND (PD<=20000327 OR PRD<=20000327 OR AD<=20000327)
                SEL DN AN 23
L46
              1 S E1-E3
              5 S L43, L46
L47
```

```
5 S L35 NOT L47
L48
L49
            369 S ?PIGR?
L50
           1490 S L8, L10, L11, L49, L13
L51
            129 S L50 AND L25
L52
             44 S L50 AND L15
L53
              8 S L51, L52 AND L19, L20
L54
              3 S L53 AND IMMUN?/SC
             2 S L54 AND LIGAND?/TI
L55
              5 S L47, L55
L56
             40 S L52 NOT L53-L56
L57
                SEL DN AN 10 15
              2 S L57 AND E4-E9
              7 S L56, L58 AND L1-L58
L59
              5 S L2-L5 AND P/DT
L60
              3 S L60 NOT L59
L61
L62
              3 S L61 AND L1-L60
             10 S L59-L62
L63
L64
              6 S L63 AND ?SECRET?
              3 S L63 AND STALK?
L65
             10 S L63-L65
L66
     FILE 'HCAPLUS' ENTERED AT 07:05:26 ON 23 JUL 2003
     FILE 'BIOSIS' ENTERED AT 07:11:59 ON 23 JUL 2003
                E MOSTOV K/AU
            199 S E3-E7
L67
                E CHAPIN S/AU
             33 S E3, E7, E9
L68
                E RICHMAN /AU
L69
             18 S E56, E60-E63
                E EISENSTAT/AU
            625 S ?PIGR?
L70
           1132 S ?POLYM? (S) IMMUNOGLOB? (S) RECEPTOR
L71
L72
             72 S L67-L69 AND L70,L71
             70 S L72 NOT PATENT/DT
L73
L74
             66 S L73 AND PY<=2000
             25 S L74 AND 00520/CC
L75
             24 S L74 AND CONFERENCE/DT
L76
L77
             25 S L75, L76
L78
             41 S L74 NOT L77
                SEL DN AN 7 21
              2 S L78 AND E1-E4
L80
              3 S L77 AND (?LIGAND? OR ANTIBOD?)
L81
             22 S L77 NOT L80
L82
             25 S L80, L81
     FILE 'BIOSIS' ENTERED AT 07:20:20 ON 23 JUL 2003
     FILE 'WPIX' ENTERED AT 07:20:45 ON 23 JUL 2003
             28 S L70/BIX
L84
            140 S L71/BIX
L85
              O S ?POLYM? (S) IMMUNO GLOB? (S) RECEPTOR/BIX
L86
            164 S L83, L84
L87
             55 S L86 AND ?LIGAND?/BIX
            125 S L86 AND ANTIBOD?/BIX
L88
L89
             31 S L87, L88 AND SECRET?/BIX
             17 S L86 AND C07K016-28/IC, ICM, ICS, ICA, ICI
L90
                SEL DN AN 5 16
              2 S L90 AND E5-E8
L91
             19 S L86 AND C07K014-705/IC, ICM, ICS, ICA, ICI
L92
L93
             11 S L92 NOT L90
                SEL DN AN 8
```

1 S L93 AND E9-E10

L94

```
36 S L86 AND A61K039-395/IC, ICM, ICS, ICA, ICI
L95
             20 S L95 NOT L90-L94
L96
L97
              3 S L91, L94
                E MOSTOV K/AU
L98
              4 S E3, E4
                E CHAPIN S/AU
              2 S E3, E5
L99
                E RICHMAN/AU
            43 S E3-E16, E20-E23
L100
             4 S L86 AND L98-L100
L101
             5 S L97, L101
L102
L103
             5 S L102 AND L83-L102
             28 S L89 NOT L103
L104
             29 S L87 NOT L89-L104
L105
     FILE 'WPIX' ENTERED AT 07:36:38 ON 23 JUL 2003
            1 S L102 NOT L101
                SEL PN L101
     FILE 'DPCI' ENTERED AT 07:39:30 ON 23 JUL 2003
              3 S E1-E19
               E MOSTOV/AU
              4 S E5, E6
L108
                E CHAPIN S/AU
L109
              1 S E5
                E RICHMAN E/AU
              3 S E3, E6
L110
L111
              5 S E9, E11
               E EISENSTAT/AU
              7 S L108-L111 NOT L107
L112
              3 S L107 AND L108-L112
L113
     FILE 'DPCI' ENTERED AT 07:41:21 ON 23 JUL 2003
     FILE 'HCAPLUS' ENTERED AT 07:49:01 ON 23 JUL 2003
L114
             4 S US5972900/PN
L115
              2 S WO9621012/PN
L116
             1 S WO9746588/PN
L117
             1 S ECKMAN ?/AU AND 1999/PY AND (21 AND 2 AND 246)/SO
L118
             1 S KRAJCI ?/AU AND 1992/PY AND (22 AND 9 AND 2309)/SO
L119
             1 S MOSTOV ?/AU AND 1994/PY AND (12 AND 63)/SO
             1 S FERKOL ?/AU AND 1993/PY AND (92 AND 5 AND 2394)/SO
L120
             6 S (US5656469 OR US5658751 OR WO9600902)/PN
L121
             1 S CARDONE ?/AU AND 1994/PY AND (124 AND 5 AND 717)/SO
L122
             1 S LIPOWSKA ?/AU AND 1993/PY AND (23 AND 21 AND 3087)/SO
L123
             2 S LEE ?/AU AND HETEROAROMAT? AND IMINIUM AND CYANIN? AND FLUORE
L124
             1 S MAZANEC ?/AU AND 1995/PY AND (69 AND 2 AND 1339)/SO
L125
              1 S WILLIAMS ?/AU AND 1988/PY AND (6 AND 2 AND 36)/SO
L126
              1 S SOLARI ?/AU AND 1985/PY AND (260 AND 1141)/SO
L127
              1 S EIFFERT ?/AU AND 1984/PY AND (365 AND 1489)/SO
L128
              1 S SOLARI ?/AU AND 1986/PY AND (34 AND 1 AND 17)/SO
L129
L130
              1 S BREITFELD ?/AU AND 1989/PY AND (109 AND 475)/SO
              1 S PISKURICH ?/AU AND 1995/PY AND (154 AND 1735)/SO
L131
              1 S FERKOL ?/AU AND 1995/PY AND (95 AND 493)/SO
L132
              1 S WU ?/AU AND 1987/PY AND (262 AND 4429)/SO
L133
             1 S BREITFELD ?/AU AND 1989/PY AND (32 AND 329)/SO
             1 S MOSTOV ?/AU AND 1994/PY AND (12 AND 63)/SO
L135
             1 S MOSTOV ?/AU AND 1984/PY AND (308 AND 5954 AND 37)/SO
L136
             1 S MOSTOV ?/AU AND 1980/PY AND (77 AND 12 AND 7257)/SO
L138
             1 S WU ?/AU AND 1987/PY AND (262 AND 4429)/SO
            0 S HUDSON ?/AU AND 1980/PY AND 192/SO
             2 S (WO200047611 OR US6207195)/PN
L140
             33 S L114-L140
L141
```

L142

31 S L141 NOT L66

FILE 'HCAPLUS' ENTERED AT 08:09:40 ON 23 JUL 2003

L143 26 S L142 AND L1-L66 L144 5 S L142 NOT L143 L145 31 S L143,L144